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Washington, DC 20549

CUTERA®

CUTERA, INC. 2011 PROXY STATEMENT AND 2010 ANNUAL REPORT

CTERA

Dear Stockholders:

You are cordially invited to attend the 2011 Annual Meeting of Stockholders of Cutera, Inc. (the "Company"). The meeting will be held at our principal executive offices located at 3240 Bayshore Blvd., Brisbane, California 94005-1021 on June 14, 2011 at 10:00 a.m. Pacific Time.

The attached Notice of 2011 Annual Meeting of Stockholders and Proxy Statement contain details of the business to be conducted at the Annual Meeting. We have also made available a copy of our 2010 Annual Report to Stockholders with this proxy statement. We encourage you to read our Annual Report. It includes our audited financial statements and provides information about our business.

We have elected to provide access to our proxy materials over the internet under the Securities and Exchange Commission's "notice and access" rules. We are constantly focused on improving the ways people connect with information, and believe that providing our proxy materials over the internet increases the ability of our stockholders to connect with the information they need, while reducing the environmental impact of our Annual Meeting. If you need additional information about Cutera, please visit the Investor Relations section of the Company's website at www.cutera.com.

Whether or not you attend the Annual Meeting, it is important that your shares be represented and voted at the meeting. Therefore, I urge you to promptly vote and submit your proxy via the internet, by phone, or by signing, dating, and returning the proxy card provided to you. If you decide to attend the Annual Meeting, you will be able to vote in person, even if you have previously submitted your proxy.

On behalf of Cutera's Board of Directors and executive team, I would like to express our appreciation for your continued interest and confidence in our business.

Sincerely,

Kevin Connors,

President and Chief Executive Officer

UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

SCHEDULE 14A

Proxy Statement Pursuant to Section 14(a) of the Securities Exchange Act of 1934

File	d by the	Securities Exchange Act of 1934 e Registrant ⊠
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File	d by a I	Party other than the Registrant □
Che	ck the a	appropriate box:
	P	reliminary Proxy Statement
	C	Confidential, for Use of the Commission Only (as permitted by Rule 14a-6(e)(2))
X	D	Definitive Proxy Statement
	Ε	Definitive Additional Materials
	S	Soliciting Material Pursuant to Sec.240.14a-11(c) or Sec.240.14a-2
		CITERA
		CUTERA, INC.
		(Name of Registrant as Specified In Its Charter)
		(Name of Person(s) Filing Proxy Statement, if other than the Registrant)
Payr	ment of	f Filing Fee (Check the appropriate box):
X	No fe	ee required.
	Fee c	computed on table below per Exchange Act Rules 14a-6(i)(4) and 0-11.
	(1)	Title of each class of securities to which transaction applies:
	(2)	Aggregate number of securities to which transaction applies:
	(3)	Per unit price or other underlying value of transaction computed pursuant to Exchange Act Rule 0-11 (Set forth the amount on which the filing fee is calculated and state how it was determined):
-	(4)	Proposed maximum aggregate value of transaction:
	(5)	Total fee paid:
	Fee p	paid previously with preliminary materials.
	offset	k box if any part of the fee is offset as provided by Exchange Act Rule 0-11(a)(2) and identify the filing for which the titing fee was paid previously. Identify the previous filing by registration statement number, or the Form or Schedule and ate of its filing.
	(1)	Amount Previously Paid:
	(2)	Form, Schedule or Registration Statement No.:
	(3)	Filing Party:
	(4)	Date Filed:

CITERA

NOTICE OF 2011 ANNUAL MEETING OF STOCKHOLDERS TO BE HELD ON JUNE 14, 2011

10:00 A.M. Pacific Time

To our Stockholders:

You are cordially invited to attend the 2011 Annual Meeting of Stockholders of Cutera, Inc. (the "Company"). The meeting will be held at our principal executive offices located at 3240 Bayshore Blvd., Brisbane, California 94005-1021. We previously disclosed in our Annual Report that the meeting will be held on June 13, 2011. The meeting, however, will be held on June 14, 2011 at 10:00 a.m. Pacific Time, for the following purposes:

- 1. To elect two Class I directors to each serve for a three-year term that expires at the 2014 Annual Meeting of Stockholders and until their successors have been duly elected and qualified;
- 2. To hold a non-binding vote on executive compensation;
- 3. To hold a non-binding vote on the frequency of executive compensation voting;
- 4. To ratify the appointment of PricewaterhouseCoopers LLP as our independent registered public accounting firm (the "Independent Registered Public Accounting Firm") for the fiscal year ending December 31, 2011; and
- 5. To transact such other business as may properly come before the Annual Meeting, including any motion to adjourn to a later date to permit further solicitation of proxies, if necessary, or before any adjournment thereof.

The foregoing items of business are more fully described in the proxy statement accompanying this Notice of Annual Meeting.

To help conserve resources and reduce printing and distribution costs, we will be mailing a notice to our stockholders, instead of a paper copy of this proxy statement and our 2010 Annual Report, with instructions on how to access our proxy materials over the Internet, including this proxy statement, our 2010 Annual Report and a form of proxy card or voting instruction card. The notice will also contain instructions on how each of those stockholders can receive a paper copy of our proxy materials.

The meeting will begin promptly at 10:00 a.m., local time, and check-in will begin at 9:50 a.m. local time. Only holders of record of shares of our common stock (NASDAQ: CUTR) at the close of business on April 18, 2011 will be entitled to notice of, and to vote at, the meeting and any postponements or adjournments of the meeting.

For a period of at least 10 days prior to the meeting, a complete list of stockholders entitled to vote at the meeting will be available and open to the examination of any stockholder for any purpose relating to the Annual Meeting during normal business hours at our principal executive offices located at 3240 Bayshore Blvd., Brisbane, California 94005-1021.

By order of the Board of Directors,

Kevin P. Connors

President and Chief Executive Officer

Brisbane, California April 26, 2011

YOUR VOTE IS IMPORTANT!

REGARDLESS OF WHETHER YOU PLAN TO ATTEND THE MEETING, PLEASE PROMPTLY VOTE BY TELEPHONE, OR IF AVAILABLE, ELECTRONICALLY, OR, IF YOU RECEIVED PER YOUR REQUEST A PAPER COPY OF OUR PROXY MATERIALS, COMPLETE, SIGN, DATE, AND RETURN THE ENCLOSED PROXY CARD IN THE ACCOMPANYING POSTAGE-PAID ENVELOPE. NO ADDITIONAL POSTAGE IS NECESSARY IF THE PROXY CARD IS MAILED IN THE UNITED STATES OR CANADA. YOU MAY REVOKE YOUR PROXY AT ANY TIME BEFORE IT IS VOTED AT THE MEETING.

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CITERA

PROXY STATEMENT FOR 2011 ANNUAL MEETING OF STOCKHOLDERS TO BE HELD ON JUNE 14, 2011

The Board of directors of Cutera, Inc., a Delaware corporation, is soliciting the enclosed proxy from you. The proxy will be used at our 2011 Annual Meeting of Stockholders to be held on Monday, June 14, 2011, beginning at 10:00 a.m., Pacific Time, which is the local time, at our principal executive offices located at 3240 Bayshore Blvd., Brisbane, California 94005-1021, and at any postponements or adjournments thereof. This proxy statement contains important information regarding the meeting. Specifically, it identifies the matters upon which you are being asked to vote, provides information that you may find useful in determining how to vote and describes the voting procedures.

In this proxy statement the terms "we", "our", "Cutera" and the "Company" each refer to Cutera, Inc.; the term "Board" means our Board of directors; the term "proxy materials" means this proxy statement, the enclosed proxy card, and our Annual Report of Form 10-K for the year ended December 31, 2010, filed with the U.S. Securities and Exchange Commission (the "SEC") on March 15, 2011, and the term "Annual Meeting" means our 2011 Annual Meeting of Stockholders.

We are sending the Notice of Internet Availability of Proxy Materials on or about May 5, 2011, to all stockholders of record at the close of business on April 18, 2011 (the "Record Date").

QUESTIONS AND ANSWERS REGARDING THIS SOLICITATION AND VOTING AT THE ANNUAL MEETING

Why am I receiving these proxy materials?

You are receiving these proxy materials from us because you were a stockholder of record at the close of business on the Record Date (which was April 18, 2011). As a stockholder of record, you are invited to attend the meeting and are entitled to and requested to vote on the items of business described in this proxy statement.

Why did I receive a notice in the mail regarding the Internet availability of the proxy materials instead of a paper copy of the proxy materials?

Pursuant to SEC rules, we have elected to provide access to our proxy materials over the Internet. Accordingly, we are sending a Notice of Internet Availability of Proxy Materials (the "Notice") to our stockholders.

All stockholders will have the ability to access the proxy materials on a website referred to in the Notice or request to receive a printed set of the proxy materials.

Instructions on how to access the proxy materials over the Internet or to request a printed copy may be found on the Notice.

In addition, stockholders may request to receive proxy materials in printed form by mail or electronically by email on an ongoing basis. Choosing to receive your future proxy materials by email will save us the cost of printing and mailing documents to you and will reduce the impact of our annual stockholders' meetings on the environment. If you chose in connection with our 2010 Annual Meeting of Stockholders to receive future proxy materials by email, you should receive an email this year with instructions containing a link to those materials and a link to the proxy voting site. In connection with our upcoming Annual Meeting, if you choose to receive future proxy materials by email, you will receive an email next year with instructions containing a link to those materials and a link to the proxy voting site. Your election to receive proxy materials by email will remain in effect until you terminate it.

What is the purpose of the Annual Meeting?

At our meeting, stockholders of record will vote upon the items of business outlined in the notice of meeting (on the cover page of this proxy statement), each of which is described more fully in this proxy statement. In addition, management will report on the performance of the Company and respond to questions from stockholders.

Who is entitled to attend the meeting?

You are entitled to attend the meeting only if you owned our common stock (or were a joint holder) as of the Record Date or if you hold a valid proxy for the meeting. You should be prepared to present photo identification for admittance.

Please also note that if you are not a stockholder of record but hold shares in street name (that is, through a broker or nominee), you will need to provide proof of beneficial ownership as of the Record Date, such as your most recent brokerage account statement, a copy of the voting instruction card provided by your broker, trustee or nominee, or other similar evidence of ownership.

The meeting will begin promptly at 10:00 a.m., local time. Check-in will begin at 9:50 a.m., local time.

Who is entitled to vote at the meeting?

Only stockholders who owned our common stock at the close of business on the Record Date are entitled to notice of and to vote at the meeting, and at any postponements or adjournments thereof.

As of the Record Date, 13,721,045 shares of our common stock were outstanding. Each outstanding share of our common stock entitles the holder to one vote on each matter considered at the meeting. Accordingly, there are a maximum of 13,721,045 votes that may be cast at the meeting.

How many shares must be present or represented to conduct business at the meeting (that is, what constitutes a quorum)?

The presence at the meeting, in person or by proxy, of the holders of a majority of the shares of our common stock entitled to vote at the meeting will constitute a quorum. A quorum is required to conduct business at the meeting. The presence of the holders of our common stock representing at least 6,860,523 votes will be required to establish a quorum at the meeting. Both abstentions and broker non-votes are counted for the purpose of determining the presence of a quorum.

What items of business will be voted on at the meeting?

The items of business scheduled to be voted on at the meeting are as follows:

- the election of two nominees to serve as Class I directors on our Board:
- 2. a non-binding vote on executive compensation;
- 3. a non-binding vote on the frequency of executive compensation voting; and
- 4. the ratification of the appointment of PricewaterhouseCoopers LLP as our Independent Registered Public Accounting Firm for the 2011 fiscal year.

These proposals are described more fully below in this proxy statement. As of the date of this proxy statement, the only business that our Board intends to present or knows of that others will present at the meeting is as set forth in this proxy statement. If any other matter or matters are properly brought before the meeting, it is the intention of the persons who hold proxies to vote the shares they represent in accordance with their best judgment.



How does the Board recommend that I vote?

Our Board recommends that you vote your shares "FOR" each of the director nominees, "FOR" the approval of a non binding vote on executive compensation and "FOR" the ratification of PricewaterhouseCoopers LLP as our Independent Registered Public Accounting Firm for the 2011 fiscal year. In addition, our Board recommends that you vote your shares for the frequency of executive compensation voting to occur every three years, rather than every year or every two years.

What shares can I vote at the meeting?

You may vote all shares owned by you as of the Record Date, including (1) shares held directly in your name as the *stockholder of record*, and (2) shares held for you as the *beneficial owner* through a broker, trustee or other nominee such as a bank.

What is the difference between holding shares as a stockholder of record and as a beneficial owner?

Most of our stockholders hold their shares through a broker or other nominee rather than directly in their own name. As summarized below, there are some distinctions between shares held of record and those owned beneficially.

Stockholders of Record. If your shares are registered directly in your name with our transfer agent, Computershare Trust Company, Inc., you are considered, with respect to those shares, the *stockholder of record*, and these proxy materials are being sent directly to you by us. As the *stockholder of record*, you have the right to grant your voting proxy directly to Cutera or to vote in person at the meeting. We have enclosed a proxy card for your use.

Beneficial Owner. If your shares are held in a brokerage account or by another nominee, you are considered the beneficial owner of shares held in street name, and these proxy materials are being forwarded to you together with a voting instruction card. As the beneficial owner, you have the right to direct your broker, trustee or nominee how to vote and are also invited to attend the meeting. Please note that since a beneficial owner is not the stockholder of record, you may not vote these shares in person at the meeting unless you obtain a "legal proxy" from the broker, trustee or nominee that holds your shares, giving you the right to vote the shares at the meeting. Your broker, trustee or nominee has enclosed or provided voting instructions for you to use in directing the broker, trustee or nominee how to vote your shares.

How can I vote my shares without attending the meeting?

Whether you hold shares directly as the stockholder of record or beneficially in street name, you may direct how your shares are voted without attending the meeting. Stockholders of record of our common stock may submit proxies by completing, signing and dating their proxy cards and mailing them in the accompanying pre-addressed envelope. Our stockholders who hold shares beneficially in street name may vote by mail by completing, signing and dating the voting instruction cards provided by the broker, trustee or nominee and mailing them in the accompanying pre-addressed envelope.

How can I vote my shares in person at the meeting?

Shares held in your name as the stockholder of record may be voted in person at the meeting. Shares held beneficially in street name may be voted in person only if you obtain a legal proxy from the broker, trustee or nominee that holds your shares giving you the right to vote the shares. Even if you plan to attend the meeting, we recommend that you also submit your proxy card or voting instructions as described above so that your vote will be counted if you later decide not to, or are unable to, attend the meeting.

Can I change my vote?

You may change your vote at any time prior to the vote at the meeting. If you are the stockholder of record, you may change your vote by granting a new proxy bearing a later date (which automatically revokes the earlier proxy), by providing a written notice of revocation to our Secretary prior to your shares being voted, or by attending the meeting and voting in person. Attendance at the meeting will not cause your previously granted proxy to be revoked unless you specifically so request.

For shares you hold beneficially in street name, you may change your vote by submitting new voting instructions to your broker, trustee or nominee, or, if you have obtained a legal proxy from your broker, trustee or nominee giving you the right to vote your shares, by attending the meeting and voting in person.

Is my vote confidential?

Proxy instructions, ballots and voting tabulations that identify individual stockholders are handled in a manner that protects your voting privacy. Your vote will not be disclosed either within Cutera or to third parties, except: (1) as necessary to meet applicable legal requirements, (2) to allow for the tabulation of votes and certification of the vote, and (3) to facilitate a successful proxy solicitation. Occasionally, stockholders provide written comments on their proxy card, which are then forwarded to our management.

What vote is required to approve each item and how are votes counted?

The vote required to approve each item of business and the method for counting votes is set forth below:

Election of Directors. The two director nominees receiving the highest number of affirmative "FOR" votes at the meeting (a plurality of votes cast) will be elected to serve as Class I directors. You may vote either "FOR" or "WITHHOLD" your vote for the director nominees. A properly executed proxy marked "WITHHOLD" with respect to the election of one or more directors will not be voted with respect to the director or directors indicated, although it will be counted for purposes of determining whether there is a quorum.

Non-binding Vote on Executive Compensation. For the non-binding vote on executive compensation, the affirmative "FOR" vote of a majority of the shares represented in person or by proxy and entitled to vote on the item will be required for approval. You may vote "FOR," "AGAINST" or "ABSTAIN" for this item of business. If you "ABSTAIN," your abstention has the same effect as a vote "AGAINST."

Non-binding Vote on the Frequency of Executive Compensation Voting. The frequency option receiving the highest number of affirmative "FOR" votes at the meeting (a plurality of votes cast) will be considered the frequency preferred by the stockholders, although such vote will not be binding on us. You may vote, with respect to the proposal to recommend, by non-binding vote, the frequency of executive compensation voting, for a vote every one, two or three years, or may abstain from voting. If you "ABSTAIN" from voting on this proposal, the abstention will not have an effect on the outcome of the vote.

Ratification of PricewaterhouseCoopers LLP as our Independent Registered Public Accounting Firm. For the ratification of the appointment of our Independent Registered Public Accounting Firm, the affirmative "FOR" vote of a majority of the shares represented in person or by proxy and entitled to vote on the item will be required for approval. You may vote "FOR," "AGAINST" or "ABSTAIN" for this item of business. If you "ABSTAIN," your abstention has the same effect as a vote "AGAINST."

What is a "broker non-vote"?

If you provide specific instructions with regard to certain items, your shares will be voted as you instruct on such items. If you sign your proxy card or voting instruction card without giving specific instructions, your shares will be voted in accordance with the recommendations of the Board ("FOR" all of the Company's nominees to the Board, "FOR" the approval, by non-binding vote, of executive compensation, "EVERY 3 YEARS" for the approval, by non-binding vote, of triennial executive compensation voting, "FOR" ratification of PricewaterhouseCoopers LLP as our Independent Registered Public Accounting Firm, and in the discretion of the proxy holders on any other matters that may properly come before the meeting).

A "broker non-vote" occurs when a broker expressly instructs on a proxy card that it is not voting on a matter, whether routine or non-routine. Under the rules that govern brokers who have record ownership of shares that are held in street name for their clients who are the beneficial owners of the shares, brokers have the discretion to vote such shares on routine matters, which includes ratifying the appointment of an independent registered public accounting firm but does not include the election of directors, the non-binding vote on executive compensation and the non-binding vote on frequency of executive compensation voting. Therefore, if you do not otherwise instruct your broker, the broker may turn in a proxy card voting your shares "FOR" ratification of the Independent Registered Public Accounting Firm. However, if you do not instruct your broker how to vote with respect to the election of directors, the non-binding vote on executive compensation and the non-binding vote on frequency of executive compensation voting, your broker may not vote with respect to such proposal and your shares will not be counted as voting in favor of these matters.

How are "broker non-votes" counted?

Broker non-votes will be counted for the purpose of determining the presence or absence of a quorum for the transaction of business, but they will not be counted in tabulating the voting result for any particular proposal.

How are abstentions counted?

If you return a proxy card that indicates an abstention from voting on all matters, the shares represented will be counted for the purpose of determining both the presence of a quorum and the total number of votes cast with respect to a proposal (other than the election of directors), but they will not be voted on any matter at the meeting. In the absence of controlling precedent to the contrary, we intend to treat abstentions in this manner. Accordingly, abstentions will have the same effect as a vote "AGAINST" a proposal.

What happens if additional matters are presented at the meeting?

Other than the two proposals described in this proxy statement, we are not aware of any other business to be acted upon at the meeting. If you grant a proxy, the persons named as proxy holders, David B. Apfelberg (the lead independent director on our Board) and Ronald J. Santilli (our Chief Financial Officer), will have the discretion to vote your shares on any additional matters that may be properly presented for a vote at the meeting. If, for any unforeseen reason, any of our nominees is not available as a candidate for director, the persons named as proxy holders will vote your proxy for such other candidate or candidates as may be nominated by our Board.

Who will serve as inspector of election?

We expect a representative of Computershare Trust Company, Inc., our transfer agent, to tabulate the votes, and expect Rajesh Madan, our Vice President of Finance to act as inspector of election at the meeting.

What should I do in the event that I receive more than one set of proxy/voting materials?

You may receive more than one set of these proxy solicitation materials, including multiple copies of this proxy statement and multiple proxy cards or voting instruction cards. For example, if you hold your shares in more than one brokerage account, you may receive a separate voting instruction card for each brokerage account in which you hold shares. In addition, If you are a stockholder of record and your shares are registered in more than one name, you may receive more than one proxy card. Please complete, sign, date and return each Cutera proxy card and voting instruction card that you receive to ensure that all your shares are voted.

Who is soliciting my vote and who will bear the costs of this solicitation?

Your vote is being solicited on behalf of the Board, and the Company will bear the entire cost of solicitation of proxies, including preparation, assembly, printing and mailing of this proxy statement. In addition to these mailed proxy materials, our directors and employees may also solicit proxies in person, by telephone, by electronic mail or by other means of communication. Directors and employees will not be paid any additional compensation for soliciting proxies. We may reimburse brokerage firms, banks and other agents for the cost of forwarding proxy materials to beneficial owners. We may also engage the services of a professional proxy solicitation firm to aid in the solicitation of proxies from certain brokers, bank nominees and other institutional owners. Our costs for such services, if retained, will not be material.

Where can I find the voting results of the meeting?

We intend to announce preliminary voting results at the Annual Meeting and file a Form 8-K with the SEC within four business days after the end of our Annual Meeting to report the voting results.

What is the deadline to propose actions for consideration at next year's Annual Meeting of stockholders or to nominate individuals to serve as directors?

As a stockholder, you may be entitled to present proposals for action at a future meeting of stockholders, including director nominations.

Stockholder Proposals: For a stockholder proposal to be considered for inclusion in our proxy statement for the Annual Meeting to be held in 2012, the written proposal must be received by our corporate Secretary at our principal executive offices no later than January 6, 2012, which is the date 120 calendar days before the anniversary of the mailing date of the Notice of Internet Availability of Proxy Materials. If the date of next year's Annual Meeting is moved more than 30 days before or after the anniversary date of this year's Annual Meeting, the deadline for inclusion of proposals in our proxy statement is instead a reasonable time before we begin to print and mail its proxy materials. Such proposals also must comply with the requirements of Rule 14a-8 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), and any other applicable rules established by the SEC. Stockholders interested in submitting such a proposal are advised to contact knowledgeable legal counsel with regard to the detailed requirements of applicable securities laws. Proposals should be addressed to:

Secretary Cutera, Inc. 3240 Bayshore Blvd. Brisbane, California 94005-1021

Nomination of Director Candidates: You may propose director candidates for consideration by our Board. Any such recommendations should include the nominee's name and qualifications for Board membership and should be directed to the "Secretary" at the address of our principal executive offices set forth above. In addition, our Bylaws permit stockholders to nominate directors for election at an Annual Meeting of stockholders. To nominate a director, the stockholder must provide the information required by our Bylaws, as well as a statement by the nominee consenting to being named as a nominee and to serve as a director if elected. In addition, the stockholder must give timely notice to our corporate Secretary in accordance with the provisions of our Bylaws, which require that the notice be received by our corporate Secretary no later than January 6, 2012.

Copy of Bylaw Provisions: You may contact our corporate Secretary at our principal executive offices for a copy of the relevant bylaw provisions regarding the requirements for making stockholder proposals and nominating director candidates.

STOCK OWNERSHIP

Security Ownership of Certain Beneficial Owners and Management

The following table provides information relating to the beneficial ownership of our common stock as of the Record Date, by:

- each stockholder known by us to own beneficially more than 5% of our common stock;
- each of our Named Executive Officers named in the Summary Compensation Table on page 23 (our Chief Executive Officer and our Chief Financial Officer);
- each of our directors; and
- all of our directors and Named Executive Officers as a group.

The number of shares beneficially owned by each entity, person, director or executive officer is determined in accordance with the rules of the SEC, and the information is not necessarily indicative of beneficial ownership for any other purpose. Under such rules, beneficial ownership includes any shares over which the individual has the sole or shared voting power or investment power and any shares that the individual has the right to acquire within 60 days of April 18, 2011 (the Record Date) through the exercise of any stock option or other right. The number and percentage of shares beneficially owned is computed on the basis of 13,721,045 shares of our common stock outstanding as of the Record Date. The information in the following table regarding the beneficial owners of more than 5% of our common stock is based upon information supplied by principal stockholders or Schedules 13D and 13G filed with the SEC.

Shares of our common stock that a person has the right to acquire within 60 days of the Record Date are deemed outstanding for purposes of computing the percentage ownership of the person holding such rights, but are not deemed outstanding for purposes of computing the percentage ownership of any other person, except with respect to the percentage ownership of all directors and executive officers as a group. To our knowledge, except as set forth in the footnotes to this table and subject to applicable community property laws, each person or entity named in the table has sole voting and disposition power with respect to the shares set forth opposite such person's or entity's name. The address for those persons for which an address is not otherwise provided is c/o Cutera, Inc., 3240 Bayshore Blvd., Brisbane, California 94005-1021.

Name and Address of Beneficial Owner	Number of Shares Outstanding	Warrants and Options Exercisable Within 60 Days	Approximate Percent Owned
Individuals and entities affiliated with Fidelity Management &	the the		
Research Company	1,284,550	_	9.4%
Individuals and entities affiliated with GAMCO Investors, Inc	1,026,285	_	7.5%
Entities affiliated with American Century Companies, Inc	835,675	_	6.1%
BlackRock, Inc	783,493		5.7%
Dimensional Fund Advisors LP	756,655		5.5%
David B. Apfelberg	26,211	52,000	*
Kevin P. Connors	558,727	278,308	6.1%
David A. Gollnick	196,165	41,126	1.7%
W. Mark Lortz	15,804	62,000	*
Timothy J. O'Shea	13,519	42,000	*
Ronald J. Santilli	34,600	178,401	1.6%
Jerry P. Widman	13,519	62,000	*
All directors and Named Executive Officers as a group (7 persons)	858,545	715,835	11.5%

^{*}Less than 1%.

Section 16(a) Beneficial Ownership Reporting Compliance

Section 16(a) of the Exchange Act requires our directors, officers and beneficial owners of more than 10% of our common stock to file reports of ownership and reports of changes in the ownership with the SEC. Such persons are required by SEC regulations to furnish us with copies of all Section 16(a) forms they file.

Based solely on our review of the copies of such forms received by us, or written representations from reporting persons that no Forms 3, 4 or 5 were required of such persons, we believe that during our fiscal year ended December 31, 2010 all reports were timely filed.

CORPORATE GOVERNANCE AND BOARD MATTERS

Director Independence

Our Board currently consists of eight authorized directors, with two vacancies. The Company's directors are David B. Apfelberg, Kevin P. Connors, David A. Gollnick, Timothy J. O'Shea, W. Mark Lortz and Jerry P. Widman. Our Board has determined that each of the directors other than Kevin P. Connors, the Company's President and Chief Executive Officer, and David A. Gollnick, the Company's former Executive Vice President of Research and Development and a current consultant to our Company satisfy the current "independent director" standards established by rules of The NASDAQ Stock Market LLC ("Nasdaq").

Board Leadership Structure

Our Board does not have a chairman. In July 2010, our Board designated David B. Apfelberg as the lead independent director of the Board. Our Chief Executive Officer, Mr. Connors, performs many of the functions that a chairman would typically perform and works together with Dr. Apfelberg in setting the agenda for each board meeting and presiding over such meetings. At the end of each board meeting, the independent directors meet without Mr. Connors and Mr. Gollnick present. Following each meeting, Dr. Apfelberg provides feedback to Mr. Connors on his performance and the performance of other Cutera employees during the meeting and frequently recommends new agenda items for the next meeting.

As described in more detail below, the Board has two standing committees, an Audit Committee and a Compensation Committee. The chairman and each member of these committees is an independent director. The Board delegates substantial duties and responsibilities to each committee. The committees make recommendations to the Board and report regularly to the Board on their activities and any actions they have taken. We believe that our independent board committees and their chairman are an important aspect of our board leadership structure.

Risk Oversight and Analysis

Our management is responsible for managing the risks we face in the ordinary course of operating our business. The Board oversees potential risks and our risk management activities by receiving operational and strategic presentations from management which include discussions of key risks to our business. While our Board has the ultimate responsibility for risk management and oversight, various committees of the Board also support the Board in its fulfillment of this responsibility. For example, our Audit Committee assists the Board in its risk oversight function by reviewing and discussing with management our system of disclosure controls and our internal controls over financial reporting, and risks associated with our cash investment policies. Our business is run conservatively and excessive risk taking has been discouraged. As a result, risk analysis has not been a significant factor for our Compensation Committee in establishing compensation.

Committees of the Board

Our Board has two standing committees: the Audit Committee and the Compensation Committee. From time to time, our Board may also create various ad hoc committees for special purposes. The membership during the last fiscal year and the function of each of the committees are described below.

Name of Director	Audit Committee	Compensation Committee
Non-Employee Directors:		
Jerry P. Widman	X*	X
Timothy J. O'Shea**	X	X
W. Mark Lortz	\mathbf{X}	
David B. Apfelberg		X*
David A. Gollnick***		- :
Employee Directors:		
Kevin P. Connors	,	
Number of Meetings Held During the Last Fiscal Year	6	4

- X = Committee member
- * = Chairman of Committee
- ** = Mr. O'Shea was appointed a member of our Compensation Committee on April 22, 2011.
- *** = Mr. Gollnick resigned from the position of Executive Vice President of Research and Development effective March 20, 2009 and continues to be a member of our Board and a consultant to our Company.

Audit Committee. The Audit Committee oversees the Company's accounting and financial reporting processes and the audits of its financial statements. In this role, the Audit Committee monitors and oversees the integrity of the Company's financial statements and related disclosures, the qualifications, independence, and performance of the Company's Independent Registered Public Accounting Firm, and the Company's compliance with applicable legal requirements and its business conduct policies. Our Board has determined that each member of the Audit Committee meets the independence and financial literacy requirements of the Nasdaq rules and the independence requirements of the SEC. Our Board has determined that Jerry P. Widman continues to qualify as an "audit committee financial expert," as defined in SEC rules. The Audit Committee has a written charter, which was adopted by our Board in January 2004, a copy of which can be found on our website at www.cutera.com. The report of the Audit Committee appears on page 14 of this proxy statement.

Compensation Committee. The Compensation Committee, together with the Board, establishes compensation for the Chief Executive Officer and the other executive officers and administers the Company's 2004 Equity Incentive Plan (as amended in 2008) and 2004 Employee Stock Purchase Plan. The Compensation Committee has a written charter, which was adopted by our Board in January 2004, and amended on April 13, 2007 and on April 25, 2008, and can be found on our website.

Meetings Attended by Directors

During 2010, the Board held five meetings, the Audit Committee held six meetings and the Compensation Committee held three meetings. No director attended fewer than 75% of the meetings of the Board or committee(s) on which he or she served during 2010.

The directors of the Company are encouraged to attend the Company's Annual Meeting of Stockholders. In 2010, director Kevin P. Connors attended the meeting in person and directors David B. Apfelberg, Timothy J. O'Shea and Jerry P. Widman attended the meeting telephonically. No other board members attended that meeting, in person or telephonically.

Director Nomination Process

Nominations. Our Board does not currently have a nominating committee or other committee performing a similar function nor do we have any formal written policies outlining the factors and process relating to the selection of nominees for consideration for Board membership by the full Board and the stockholders. Our Board has adopted resolutions in accordance with the Nasdaq Rules authorizing a majority of its independent members to recommend qualified nominees for consideration by the full Board. Our Board believes that it is appropriate for us to not have a standing nominating committee because of a number of factors, including the number of independent directors who want to participate in consideration of candidates for membership on the Board. Our Board consists of six members, four of whom are independent. Our Board considered forming a nominating committee consisting of several of the independent members of our Board. Forming a committee consisting of less than all of the independent members would have resulted in the omission of the other independent members of our Board who wanted to participate in considering qualified candidates for Board membership. Since our Board desired the participation in the nominations process of all of its independent members, it therefore decided not to form a nominating committee and instead authorized a majority of the independent members of our Board to make and consider nominations for Board membership. The independent members of our Board do not have a nominating committee charter, but act pursuant to Board resolutions as described above and according to written nominating and corporate governance guidelines adopted in October 2010. Each of the members of our Board authorized to recommend nominees to the full Board is independent within the meaning of the current "independent director" standards established by Nasdaq's rules. Our Board last reviewed this matter in October 2010 and intends to review it periodically, and may in the future elect to designate a formal nominating committee.

Director Qualifications. While the independent members of our Board have not established specific minimum qualifications for director candidates, the candidates for Board membership should have the highest professional and personal ethics and values, and conduct themselves consistent with our Code of Ethics. While the independent members of the Board have not formalized specific minimum qualifications they believe must be met by a candidate to be recommended by the independent members, the independent members of the Board believe that candidates and nominees must reflect a Board that is comprised of directors who (i) have broad and relevant experience, (ii) are predominantly independent, (iii) are of high integrity, (iv) have qualifications that will increase overall Board effectiveness and enhance long-term stockholder value, and (v) meet other requirements as may be required by applicable rules, such as financial literacy or financial expertise with respect to Audit Committee members.

Stockholder Nominations and Recommendations. As described above in the Question and Answer section of this proxy statement under "What is the deadline to propose actions for consideration at next year's Annual Meeting of Stockholders or to nominate individuals to serve as directors?," our Bylaws set forth the procedure for the proper submission of stockholder nominations for membership on our Board. In addition, the independent members of our Board may consider properly submitted stockholder recommendations (as opposed to formal nominations) for candidates for membership on the Board. A stockholder may make such a recommendation by submitting the following information to our Secretary at 3240 Bayshore Blvd., Brisbane, California 94005-1021: the candidate's name, home and business contact information, detailed biographical data, relevant qualifications, professional and personal references, information regarding any relationships between the candidate and Cutera within the last three years and evidence of ownership of Cutera stock by the recommending stockholder.

Identifying and Evaluating Director Nominees. Typically new candidates for nomination to the Board are suggested by existing directors or by our executive officers, although candidates may initially come to our attention through professional search firms, stockholders or other persons. The independent members of the Board carefully review the qualifications of any candidates who have been properly brought to their attention. Such a review may, in the Board's discretion, include a review solely of information provided to the Board or may also include discussion with persons familiar with the candidate, an interview with the candidate or other actions that the Board deems proper. The Board shall consider the suitability of each candidate, including the current members of the Board, in light of the current size and composition of the Board. In evaluating the qualifications of the candidates, the independent members of the Board considers many factors, including, issues of character, judgment, independence, expertise, length of service, and other commitments. In addition, the independent members of the Board take into account diversity in professional experience, skills and background in considering and evaluating candidates. However, while diversity relating to background, skill, experience and perspective is one factor considered in the nomination process, the Company does not have a formal policy relating to diversity. The Board evaluates such factors, among others, and does not assign any particular weighting or priority to any of these factors. Candidates properly recommended by stockholders are evaluated by the independent directors using the same criteria as other candidates. Candidates are not discriminated against on the basis of race, religion, national origin, sexual orientation, disability or any other basis proscribed by law.

Director Compensation

The following table sets forth a summary of the cash compensation and the grant date fair value of fully vested Cutera stock awarded to our non-employee directors in the year ended December 31, 2010.

Name	0	es Earned r Paid in Cash ⁽¹⁾	Stock Awards (2)	Comp	l Other ensation (3)	Total
David B. Apfelberg	\$	65,000	\$ 60,000 ⁽⁴⁾	\$		\$ 125,000
David A. Gollnick		45,000	$60,000^{(5)}$		$133,520^{(5)}$	238,520
W. Mark Lortz		52,500	$60,000^{(6)}$		·	112,500
Timothy J. O'Shea		52,500	$60,000^{(7)}$		· . 	112,500
Jerry P. Widman		71,000	$60,000^{(8)}$			131,000

- (1) Amounts were earned in connection with serving on our Board and its committees, or committee Chairman retainers, each as described below.
- Amounts shown in this column are the aggregate grant date fair value of fully vested stock awards granted during the year ended December 31, 2010 calculated in accordance with Accounting Standards Codification (ASC) Topic 718.
- (3) Amounts shown in this column were earned outside of serving on the Board or its committees or are reportable and do not meet the definition of fees earned or paid in cash or stock awards, each as described below.
- (4) At December 31, 2010, David B. Apfelberg held options to purchase 52,000 shares of common stock.
- David A. Gollnick resigned from the position of Executive Vice President of Research and Development effective March 20, 2009. He continues to be a member of our Board and is a consultant to our Company. In connection with his consulting agreement, he was paid \$133,250 during the year ended December 31, 2010. At December 31, 2010, Mr. Gollnick held options to purchase 41,126 shares of common stock.
- (6) At December 31, 2010, W. Mark Lortz held options to purchase 62,000 shares of common stock.
- (7) At December 31, 2010, Timothy J. O'Shea held options to purchase 42,000 shares of common stock.
- (8) At December 31, 2010, Jerry P. Widman held options to purchase 62,000 shares of common stock.

From January 1, 2009 through June 28, 2009, our non-employee directors earned an annual retainer of \$25,000 for regular board meetings. In June 2009, the Board reviewed a report from Mercer, the Company's compensation consultant, relating to director compensation. The results of the Mercer report demonstrated that:

- there was a trend away from per meeting fees and towards annual retainers for services rendered by Board and Committee members;
- 2) cash compensation for directors was increasing based on the additional duties and responsibilities of Board and committee members; and
- 3) there was a trend away from stock options to fully vested equity grants based on a fixed dollar value rather than a share value.

For 2010, our non-employee directors earned an annual retainer of \$45,000 for regular board meetings; \$6,000 for Compensation Committee meetings (for members other than the Chairman); and \$7,500 for Audit Committee meetings (for members other than the Chairman). The Chairman of the Audit Committee and the Chairman of the Compensation Committee, each earned an annual retainer of \$20,000 for their services on the respective committees. Our directors no longer receive per meeting fees for board and committee meetings regardless of the number of meetings held throughout the year.

Our 2004 Equity Incentive Plan provides for the automatic grant of options to our non-employee directors. Each non-employee director who is appointed to the Board will receive an initial option to purchase 14,000 shares of our common stock upon such appointment. Each option to purchase 14,000 shares will have an exercise price equal to fair market value on the date of grant and a term of seven years and will become exercisable as to one-third of the shares subject to the option on each anniversary of its date of grant, provided the non-employee director remains a director on such dates. In addition, each non-employee director who is a director on the date of each Annual Meeting of stockholders and has been a director for at least the preceding six months, will receive fully vested Cutera stock on an annual basis equivalent to the number of shares represented by the quotient of \$60,000 divided by the closing stock price of our common stock on the date of such Annual Meeting. In May 2010, our non-employee directors received \$60,000 of fully vested Cutera stock.

Code of Ethics

We are committed to maintaining the highest standards of business conduct and ethics. Our Code of Ethics, as amended, (the "Code") reflects our values and the business practices and principles of behavior that support this commitment. The Code is intended to satisfy SEC rules for a "code of ethics" required by Section 406 of the Sarbanes-Oxley Act of 2002, as well as the Nasdaq listing standards requirement for a "code of conduct." The Code is an Exhibit to our Form 8-K filed with the SEC on April 27, 2004, was amended and restated on November 19, 2009, and is available on the Company's website at www.cutera.com. We will post any amendment to the Code, as well as any waivers that are required to be disclosed by the rules of the SEC or Nasdaq, on our website.

Corporate Governance Committee Guidelines

We do not have a Corporate Governance Committee but our Board adopted a corporate governance policy and corporate governance guidelines in October 2010. Our Board, together with our committees, fulfills the role that a Corporate Governance Committee would provide and work with management on corporate governance matters generally.

Compensation Committee Interlocks and Insider Participation

No member of our Compensation Committee, nor any of our executive officers, has a relationship that would constitute an interlocking relationship with executive officers or directors of another entity. No Compensation Committee member is an officer or employee of Cutera.

Certain Relationships and Related Transactions

In the Company's last fiscal year, and except for compensation paid to its directors and executive officers for services performed in such roles, and except as provided in the following paragraph, there has not been, nor is there currently proposed, any transaction or series of similar transactions to which the Company was or is to be a party in which the amount involved exceeds \$120,000 and in which any director, executive officer, holder of more than 5% of our common stock or any member of their immediate families had or will have a direct or indirect material interest.

We have a consulting agreement with David A. Gollnick pursuant to which Mr. Gollnick is compensated for services that he provides to us, including product development and clinical support. Payments to Mr. Gollnick under this agreement in 2010 and 2009 were \$133,520 and \$110,400, respectively.

Review, Approval or Ratification of Related Party Transactions

As provided by our Audit Committee charter, our Audit Committee must review and approve in advance any proposed related party transaction. All of our directors and officers are required to report to our Audit Committee any such related party transaction prior to its completion. We have not adopted specific standards for approval of related party transactions, but instead our Audit Committee reviews each such transaction on a case-by-case basis. Our policy is to require that all executive compensation-related matters be recommended and approved by our Compensation Committee as provided by our Compensation Committee charter and be reported under applicable SEC rules.

Family Relationships

There are no family relationships among any of our directors or executive officers.

Indemnification Agreements

Each of our directors and officers has an indemnification agreement with our Company.

Communications with the Board by Stockholders

Stockholders wishing to communicate with the Board or with an individual Board member concerning the Company may do so by writing to the Board or to the particular Board member, and mailing the correspondence to: Attention: Board of Directors, c/o Secretary, Cutera, Inc., 3240 Bayshore Blvd., Brisbane, California 94005-1021. The envelope should indicate that it contains a stockholder communication. All such stockholder communications will be forwarded to the director or directors to whom the communications are addressed.

REPORT OF THE AUDIT COMMITTEE

The material in this section is not deemed filed with the SEC and is not incorporated by reference in any filing of our Company under the Securities Act of 1933 or the Securities Exchange Act of 1934, whether made before or after the date of this Proxy Statement and irrespective of any general incorporation language in those filings.

The Audit Committee of the Board of Directors is comprised solely of independent directors (as defined by Nasdaq rules) who were all appointed by the Board of Directors. The Audit Committee operates pursuant to a written charter adopted by the Board of Directors, a copy of which can be found on our website. The Audit Committee reviews and assesses the adequacy of its charter on an annual basis. As more fully described in the charter, the purpose of the Audit Committee is to provide general oversight of Cutera's financial reporting, integrity of financial statements, internal controls and internal audit functions. The Audit Committee has authority to retain outside legal, accounting or other advisors as its deems necessary to carry out its duties and to require Cutera to pay for such expenditures.

The Audit Committee monitors Cutera's external audit process, including the scope, fees, auditor independence matters and the extent to which the Independent Registered Public Accounting Firm may be retained to perform non-audit services. The Audit Committee has responsibility for the appointment, compensation, retention and oversight of Cutera's Independent Registered Public Accounting Firm. The Audit Committee also reviews the results of the external audit work with regard to the adequacy and appropriateness of Cutera's financial, accounting and internal controls over financial reporting. In addition, the Audit Committee generally oversees Cutera's internal compliance programs. The Audit Committee members are not all professional accountants or auditors, and their function is not intended to duplicate or to certify the activities of management and the Independent Registered Public Accounting Firm, nor can the Audit Committee certify that the Independent Registered Public Accounting Firm is "independent" under applicable rules.

The Audit Committee provides advice, counsel and direction to management and the Independent Registered Public Accounting Firm on matters for which it is responsible based on the information it receives from management and the Independent Registered Public Accounting Firm and the experience of its members in business, financial and accounting matters.

Management is responsible for the preparation and integrity of Cutera's financial statements, accounting and financial reporting processes and internal control over financial reporting for compliance with applicable accounting standards, laws and regulations.

Cutera's Independent Registered Public Accounting Firm, PricewaterhouseCoopers LLP, is responsible for performing an independent audit of Cutera's financial statements in accordance with generally accepted auditing standards and expressing an opinion in its report on those financial statements, and for expressing an opinion on the effectiveness of Cutera's internal control over financial reporting.

In this context, the Audit Committee hereby reports as follows:

- The Audit Committee has reviewed and discussed the audited financial statements for 2010 with Cutera's management.
- The Audit Committee has discussed with the Independent Registered Public Accounting Firm the matters required to be discussed by SAS 61 (Codification of Statements on Auditing Standard, AU 380), SAS 99 (Consideration of Fraud in a Financial Statement Audit) and SEC rules discussed in Final Releases Nos. 33-8183 and 33-8183a.

The Audit Committee has received written disclosures and a letter from the Independent Registered Public Accounting Firm, PricewaterhouseCoopers LLP, pursuant to Rule 3526, Communication with Audit Committees Concerning Independence, of the Public Company Accounting Oversight Board ("PCAOB"), and has discussed with PricewaterhouseCoopers LLP its independence.

- The Audit Committee has discussed with the Independent Registered Public Accounting Firm the overall scope and plans for its audit.
- The Audit Committee has met with the Independent Registered Public Accounting Firm, with and
 without management present, to discuss the results of its examinations, its evaluations of our
 internal control over financial reporting, and to discuss the overall quality of our financial
 reporting.
- The Audit Committee has considered whether the provision by the Independent Registered Public Accounting Firm of non-audit services is compatible with maintaining its independence.
- Based on the review and discussion referred to above, the Audit Committee has approved that the audited financial statements and the report of management on internal control over financial reporting be included in Cutera's Annual Report on Form 10-K for the fiscal year ended December 31, 2010.

The foregoing report is provided by the undersigned members of the Audit Committee.

W. Mark Lortz Timothy J. O'Shea Jerry P. Widman

PROPOSAL ONE—ELECTION OF DIRECTORS

Classes of the Board of Directors

Our Amended and Restated Certificate of Incorporation provides that our Board shall be divided into three classes designated as Class I, Class II and Class III, respectively, with the classes of directors serving for staggered three-year terms. Our Board currently consists of six directors, divided among the three classes as follows:

- two Class I directors, Kevin P. Connors and David A. Gollnick, whose terms expire at our Annual Meeting of Stockholder to be held in 2011;
- two Class II directors, David B. Apfelberg and Timothy J. O'Shea, whose terms expire at our Annual Meeting of Stockholders to be held in 2012; and
- two Class III directors W. Mark Lortz and Jerry P. Widman, whose terms expire at the Annual Meeting of Stockholders to be held in 2013.

The name of each member of the Board, the class in which he or she serves, and his or her age as of the Record Date, principal occupation and length of service on the Board are as follows:

Name	Term Expires	Age	Principal Occupation	Director Since
Class I Directors		_		
Kevin P. Connors	2011	49	President and Chief Executive Officer	1998
David A. Gollnick	2011	47	Former Executive Vice President of Research and Development	1998
Class II Directors				:
Timothy J. O'Shea(1)(2)	2012	58	Managing Director, Oxo Capital	2004
David B. Apfelberg(1)	2012	69	Clinical Professor of Plastic Surgery, Stanford University Medical Center	1998
Class III Directors				
W. Mark Lortz (2)	2013	59	Former Chief Executive Officer, TheraSense, Inc.	2004
Jerry P. Widman (1)(2)	2013	68	Former Chief Financial Officer, Ascension Health	2004

⁽¹⁾ Member of the Compensation Committee.

Director Nominees

The Board has nominated Kevin P. Connors and David A. Gollnick for re-election as Class I directors.

Kevin P. Connors has served as our President and Chief Executive Officer and as a member of our board of directors since our inception in August 1998. From May 1996 to June 1998, Mr. Connors served as President and General Manager of Coherent Medical Group, a unit of Coherent Inc., which manufactures lasers, optics and related accessories. We believe Mr. Connors' qualifications to serve on our board of directors include, his knowledge of and leadership experience, in the aesthetic medical equipment industry prior to joining Cutera and the substantial understanding of the Company and its operations that he has gained while serving as President, Chief Executive Officer and director of the Company since inception.

⁽²⁾ Member of the Audit Committee.

David A. Gollnick has served as a member of our Board since our inception in August 1998. He served as our Vice President of Research and Development from August 1998 until April 2007, and served as our Executive Vice President of Research and Development from April 2007 until March 2009. From June 1996 to July 1998, Mr. Gollnick was Vice President of Research and Development at Coherent Medical Group, a unit of Coherent Inc. Mr. Gollnick holds a B.S. in Mechanical Engineering from Fresno State University. We believe Mr. Gollnick's qualifications to serve on our board of directors include his technical experience in researching and developing products for the aesthetic medical equipment industry and his understanding of our employees, products and operations.

If elected to our board of directors, directors Kevin P. Connors and David A. Gollnick would each hold office as a Class I director until our Annual Meeting of Stockholders to be held in 2014 or until his earlier resignation, removal, or death.

Board of Directors' Recommendation

THE BOARD OF DIRECTORS RECOMMENDS THAT YOU VOTE "FOR" EACH OF THE TWO NOMINEES FOR CLASS I DIRECTOR LISTED ABOVE.

Directors Whose Terms Extend Beyond the 2011 Annual Meeting

David B. Apfelberg, MD has served as a member of our board of directors since November 1998. Since 1980, Dr. Apfelberg has held various roles at the Stanford University Medical Center, and currently serves as a Clinical Professor of Plastic Surgery. Since 1987, Dr. Apfelberg has also been a consultant for entrepreneurs and venture capital companies in the areas of medical devices and medicine. From June 1991 to May 2001, Dr. Apfelberg was Director of the Plastic Surgery Center in Atherton, California. Dr. Apfelberg holds both a B.M.S., Bachelor of Medical Science, and an M.D. from Northwestern University Medical School. We believe Dr. Apfelberg's qualifications to serve on our board of directors include his medical expertise, understanding of our products, and his knowledge of the aesthetics market generally.

Timothy J. O'Shea has served as a member of our board of directors since April 2004. Mr. O'Shea has been with Oxo Capital since 2008 and serves as a managing director. From 1995 to 2008, he served in a variety of management positions at Boston Scientific, including Corporate Vice President of Business Development from 2000 to 2008. Mr. O'Shea holds a B.A. in history from the University of Detroit. We believe Mr. O'Shea's qualifications to serve on our board of directors include his corporate marketing knowledge as well as his diverse experience in the medical device industry working for a large medical device company.

W. Mark Lortz has served as a member of our board of directors since June 2004. Mr. Lortz served as the Chairman, President and Chief Executive Officer of TheraSense until June of 2004 after its acquisition by Abbott Laboratories. Prior to TheraSense, Mr. Lortz held several positions at LifeScan, including Vice President, Operations and Group Vice President, Worldwide Business Operations. Prior to LifeScan, Mr. Lortz had 18 years of experience with the General Electric Company in several divisions. Mr. Lortz currently serves as a member of the board of directors of two privately-held companies in the healthcare industry. Within the past five years, Mr. Lortz also served on the board of directors of NeuroMetrix, a publicly-traded manufacturer of neurological diagnostic and therapeutic devices, and IntraLase, a manufacturer of lasers for the medical industry and for eye surgery, which was acquired by Advanced Medical Optics. Mr. Lortz holds an M.B.A. in Management from Xavier University and a B.S. in Engineering Science from Iowa State University. We believe Mr. Lortz's qualifications to serve on our board of directors include his executive leadership and management experience as a former Chief Executive Officer, as well as his experience serving on the boards of other public and private companies.

Jerry P. Widman has served as a member of our board of directors since March 2004. From 1982 to 2001, Mr. Widman served as the Chief Financial Officer of Ascension Health, a not-for-profit multi-hospital system. Mr. Widman currently serves as a member of the board of directors of three other privately-held companies in the healthcare industry. Within the past five years, Mr. Widman also served on the board of directors of ArthroCare Corporation, United Surgical Partners International and the Trizetto Group. Mr. Widman holds a B.B.A. from Case Western Reserve University, an M.B.A. from the University of Denver, and a J.D. from Cleveland State University and is a Certified Public Accountant. We believe Mr. Widman's qualifications to serve on our board of directors include his financial expertise and prior experience as a Chief Financial Officer, as well as his experience serving on the boards of various public and private companies.

PROPOSAL TWO—NON-BINDING VOTE ON EXECUTIVE COMPENSATION

General

The recently enacted Dodd-Frank Wall Street Reform and Consumer Protection Act of 2010 (the "Dodd-Frank Act") enables our stockholders to vote to approve, on an advisory or non-binding basis, the compensation of our Named Executive Officers as disclosed in accordance with the SEC's rules in the "Executive Compensation" section of this proxy statement beginning on page 23 below. This proposal, commonly known as a "say-on-pay" proposal, gives our stockholders the opportunity to express their views on our Named Executive Officers' compensation as a whole. This vote is not intended to address any specific item of compensation or any specific named executive officer, but rather the overall compensation of all of our Named Executive Officers and the philosophy, policies and practices described in this proxy statement.

The say-on-pay vote is advisory, and therefore not binding on the Company, the Compensation Committee or our Board. The say-on-pay vote will, however, provide information to us regarding investor sentiment about our executive compensation philosophy, policies and practices, which the Compensation Committee will be able to consider when determining executive compensation for the remainder of the current fiscal year and beyond. Our Board and our Compensation Committee value the opinions of our stockholders and to the extent there is any significant vote against the named executive officer compensation as disclosed in this proxy statement, we will consider our stockholders' concerns and the Compensation Committee will evaluate whether any actions are necessary to address those concerns.

Summary of 2010 Executive Compensation Program

Following is a summary of some of the key points of our 2010 executive compensation program:

- The primary objectives of our compensation programs are that they be fair, objective and consistent across the employee population, that compensation be directly and substantially linked to measurable corporate and individual performance and that compensation remains competitive, so that we can attract, motivate, retain and reward the key employees whose knowledge, skills and performance are necessary for our success.
- •. We seek to foster a culture where individual performance is aligned with organizational objectives.
- We evaluate and reward our Named Executive Officers based on the comparable industry specific and general market compensation for their respective positions in the company and an evaluation of their contributions to the achievement of short-and long-term organizational goals.
- Executive compensation is reviewed annually by our Compensation Committee, and adjustments are made to reflect performance-based factors and competitive conditions.
- Our Named Executive Officers are compensated with cash, equity and non-equity incentives, and other customary employee benefits.
- Our Named Executive Officers' have Change of Control and Severance Agreements and except for these Change of Control and Severance Agreements, we do not have employment agreements with any of our Named Executive Officers.

See the "Executive Compensation" section beginning on page 23 below for more information.

Cutera believes that the information provided above and within the Executive Compensation section of this proxy statement demonstrates that it's executive compensation program was designed appropriately and is working to ensure management's interests are aligned with our stockholders' interests to support long-term value creation.

Accordingly, we ask our stockholders to vote "FOR" the following resolution at the Annual Meeting:

"RESOLVED, that the Company's stockholders approve, on an advisory basis, the compensation of the Named Executive Officers, as disclosed in the Company's Proxy Statement for the Annual Meeting of Stockholders pursuant to the compensation disclosure rules of the Securities and Exchange Commission, including the Compensation Discussion and Analysis, the compensation tables and the other related disclosure."

Board of Directors' Recommendation

THE BOARD OF DIRECTORS RECOMMENDS THAT YOU VOTE "FOR" THE ADVISORY (NON-BINDING) VOTE APPROVING EXECUTIVE COMPENSATION.

PROPOSAL THREE—NON-BINDING VOTE ON THE FREQUENCY OF EXECUTIVE COMPENSATION VOTING

The Dodd-Frank Act also enables our stockholders to indicate, at least once every six years, how frequently we should seek a non-binding vote on the compensation of our Named Executive Officers, as disclosed pursuant to the SEC's compensation disclosure rules, such as Proposal Two beginning on page 18 of this proxy statement. By voting on this Proposal Three, stockholders may indicate whether they would prefer a non-binding vote on Named Executive Officer compensation once every one, two, or three years.

After careful consideration, our Board of Directors has determined that a non-binding vote on executive compensation that occurs triennially is appropriate for the Company, and therefore our Board of Directors recommends that you vote for a three-year period between the non-binding vote on executive compensation.

In formulating its recommendation, our Board of Directors considered that given the nature of our compensation programs, a triennial vote would be sufficient for our stockholders to provide us with their input on our compensation philosophy, policies and practices. Our compensation programs do not change significantly from year-to-year and do not contain any significant risks that we believe would be of concern to our stockholders. A triennial approach provides regular input by stockholders, while allowing time to evaluate the effects of our compensation program on performance over a longer period. We understand that our stockholders may have different views as to what is the best approach in this regard, and we look forward to hearing from our stockholders on this Proposal.

You may cast your vote on your preferred voting frequency by choosing the option of one year, two years, three years or abstain from voting when you vote in response to the resolution set forth below:

"RESOLVED, that the option of once every one year, two years, or three years that receives the highest number of votes cast for this resolution will be determined to be the preferred frequency with which the Company is to hold a stockholder vote to approve the compensation of the Named Executive Officers, as disclosed pursuant to the compensation disclosure rules of the Securities and Exchange Commission, including the Compensation Discussion and Analysis, the compensation tables and the other related disclosure."

The option of one year, two years or three years that receives the highest number of votes cast by stockholders will be the frequency for the advisory vote on executive compensation that has been selected by stockholders. However, because this vote is advisory and not binding on the Company, the Compensation Committee or our Board may decide that it is in the best interests of our stockholders and the Company to hold an advisory vote on executive compensation more or less frequently than the option approved by our stockholders.

Board of Directors' Recommendation

THE BOARD OF DIRECTORS RECOMMENDS A TRIENNIAL VOTE AS THE FREQUENCY WITH WHICH STOCKHOLDERS ARE PROVIDED AN ADVISORY (NON-BINDING) VOTE ON EXECUTIVE COMPENSATION.

PROPOSAL FOUR—RATIFICATION OF PRICEWATERHOUSECOOPERS LLP AS OUR INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM

The Audit Committee of the Board has selected PricewaterhouseCoopers LLP as the Independent Registered Public Accounting Firm to perform the audit of the Company's consolidated financial statements for the fiscal year ending December 31, 2011. PricewaterhouseCoopers LLP audited the Company's consolidated financial statements for the fiscal years 2001 through 2010.

The Board is asking the stockholders to ratify the selection of PricewaterhouseCoopers LLP as the Company's Independent Registered Public Accounting Firm for 2011. Although not required by law, by rules of NASDAQ, or by the Company's bylaws, the Board is submitting the selection of PricewaterhouseCoopers LLP to the stockholders for ratification as a matter of good corporate practice. Even if the selection is ratified, the Audit Committee in its discretion may select a different Independent Registered Public Accounting Firm at any time during the year if it determines that such a change would be in the best interests of the Company and its stockholders.

Representatives of PricewaterhouseCoopers LLP are expected to be present at the Annual Meeting. They will have an opportunity to make a statement if they desire to do so and will be available to respond to appropriate questions from the Company's stockholders.

Board of Directors' Recommendation

THE BOARD OF DIRECTORS RECOMMENDS THAT YOU VOTE FOR THE RATIFICATION OF THE SELECTION OF PRICEWATERHOUSECOOPERS LLP AS THE COMPANY'S INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM FOR 2011.

Audit and Non-Audit Services

The Audit Committee is directly responsible for the appointment, compensation, and oversight of the Company's Independent Registered Public Accounting Firm. In addition to retaining PricewaterhouseCoopers LLP to audit the Company's consolidated financial statements for 2010, the Audit Committee retained PricewaterhouseCoopers LLP to provide other auditing and advisory services in 2010. The Audit Committee understands the need for PricewaterhouseCoopers LLP to maintain objectivity and independence in its audits of the Company's financial statements. The Audit Committee has reviewed all non-audit services provided by PricewaterhouseCoopers LLP in 2010 and has concluded that the provision of such services was compatible with maintaining PricewaterhouseCoopers LLP's independence in the conduct of its auditing functions.

To help ensure the independence of the Independent Registered Public Accounting Firm, the Audit Committee has adopted a policy for the pre-approval of all audit and non-audit services to be performed for the Company by its Independent Registered Public Accounting Firm. Pursuant to this policy, all audit and non-audit services to be performed by the Independent Registered Public Accounting Firm must be approved in advance by the Audit Committee. The Audit Committee may delegate to one or more of its members the authority to grant the required approvals, provided that any exercise of such authority is presented to the full Audit Committee at its next regularly scheduled meeting.

All of the services provided by PricewaterhouseCoopers LLP described in the table below were approved by the Audit Committee.

The aggregate fees incurred by the Company for audit and non-audit services in 2010 and 2009 were as follows:

Service Category	2010	2009 ⁽⁵⁾
Audit Fees ⁽¹⁾	\$ 507,150	\$ 514,750
Audit Related Fees ⁽²⁾	11.200	
Tax Fees ⁽³⁾	102,900	_
All Other Fees ⁽⁴⁾	1,500	1,500
Total	\$ 622,750	\$ 516,250

- (1) In accordance with the SEC's definitions and rules, audit fees are comprised of billed and unbilled fees for professional services related to the audit of financial statements and internal control over financial reporting for the Company's 2010 and 2009 fiscal years as included in the annual report on Form 10-K; and the review of financial statements for interim periods included in the quarterly reports on Form 10-Q within those years.
- (2) Audit-related fees are fees for services that are normally provided by the accountant in connection with statutory and regulatory filings or engagements;
- (3) Tax fees are fees for tax compliance services;
- (4) All other fees relates to a subscription fee for a PricewaterhouseCoopers LLP online service used for accounting research purposes.
- (5) The company revised the audit and non-audit fee disclosure amounts for 2009 to present data that is consistent with the 2010 disclosure, which was based on fees billed and unbilled for the fiscal years 2010 and 2009. The data presented in the prior year's proxy was based on fees incurred during the respective years.

NAMED EXECUTIVE OFFICERS AND EXECUTIVE COMPENSATION

Set forth below is certain information as of the Record Date concerning our Named Executive Officers who were with the Company as of December 31, 2010.

Name	Age	Position(s)
Kevin P. Connors	49	President, Chief Executive Officer and Director
Ronald J. Santilli	51	Executive Vice President and Chief Financial Officer

Further information regarding Kevin P. Connors is provided above under "Director Nominees."

Ronald J. Santilli has served as our Chief Financial Officer since September 2001 and as our Executive Vice President since April 2007. From September 2001 to April 2007, Mr. Santilli served as our Vice President of Finance and Administration. From April 2001 to August 2001, Mr. Santilli served as Senior Director of Financial Planning and Accounting at Lumenis, a manufacturer of medical lasers. From May 1993 to March 2001, Mr. Santilli held several positions at Coherent Inc., including Sales Operations Manager, Controller of the Medical Group and, most recently, Director of Finance and Administration. Mr. Santilli holds a B.S. in Business Administration from San Jose State University and an M.B.A. in Finance from Golden Gate University.

Compensation Discussion and Analysis

Overview

The primary objectives of our compensation programs are:

- that they be fair, objective and consistent across the employee population;
- that compensation be directly and substantially linked to measurable corporate and individual performance; and
- that compensation remains competitive, so that we can attract, motivate, retain and reward the key employees whose knowledge, skills and performance are necessary for our success.

We seek to foster a culture where individual performance is aligned with organizational objectives. We evaluate and reward our Named Executive Officers based on the comparable industry specific and general market compensation for their respective positions in the company and an evaluation of their contributions to the achievement of short-and long-term organizational goals. Executive compensation is reviewed annually by our Compensation Committee, and adjustments are made to reflect performance-based factors and competitive conditions.

Role of Our Compensation Committee

Compensation Committee Charter

The Compensation Committee establishes compensation for our two Named Executive Officers – our Chief Executive Officer and Chief Financial Officer, and administers our equity incentive plans, which are currently the 2004 Equity Incentive Plan and the 2004 Employee Stock Purchase Plan. The Compensation Committee has a written charter, which was adopted by our Board in January 2004, and was amended in April 2007 and in April 2008. A copy of this charter, as amended, can be found on our website, which is www.cutera.com.

Duties of the Compensation Committee

The responsibilities of the Compensation Committee include:

- (i) Establishing the following for the Named Executive Officers and such other officers as appropriate: (a) annual base salary, (b) annual incentive bonus, which may include the setting of specific goals and amounts, (c) equity compensation, (d) agreements for employment, severance and change-of-control, and (e) any other benefits, compensation or arrangements, other than benefits generally available to our employees.
- (ii) Reviewing and making recommendations to our Board of Directors, at such intervals as may be decided by the Compensation Committee from time to time, regarding (a) general compensation goals and guidelines for our employees and the criteria by which bonuses and stock compensation awards to our employees are determined; and, (b) other policies and plans for the provision of compensation to our employees, directors and consultants.
- (iii) Acting as Administrator of our 2004 Equity Incentive Plan, 2004 Employee Stock Purchase Plan, and any other equity compensation plans adopted by our Board.
- (iv) Reviewing and making recommendations to our Board with respect to policies relating to the issuance of equity incentives to employees, consultants and directors.
 - (v) Evaluating the compensation of the independent members of our Board.
 - (vi) Preparing the report that follows this Compensation Discussion and Analysis.

Compensation Committee Members

The members of our Compensation Committee are appointed by our Board. The members of that committee as of the Record Date were Dr. David B. Apfelberg (chairman) and Mr. Jerry P. Widman. On April 22, 2011, the Board appointed Timothy J. O'Shea as a member of the Compensation Committee. Each member of the Compensation Committee is an "outside director" for purposes of Section 162(m) of the Internal Revenue Code, a "non-employee director" for purposes of Rule 16b-3 under the Exchange Act and satisfies the independence requirements imposed by Nasdaq.

Role of the Compensation Committee and its Consultant in Setting Executive Compensation

Our Compensation Committee establishes the compensation packages for our Named Executive Officers to ensure consistency with market compensation rates for similar positions, our compensation philosophy and corporate governance guidelines. Decisions are made only by the directors who are "outside directors" for purposes of Section 162(m) of the Internal Revenue Code and "non-employee directors" for purposes of Rule 16b-3 under the Exchange Act.

With the SEC's recent reforms relating to executive compensation disclosure, our Compensation Committee has assumed an active role in reviewing market data and working with a compensation consultant on executive compensation matters. We worked with a third-party compensation consultant in March 2009, Mercer, to assist us in establishing executive compensation. Because certain components of executive compensation—such as bonus targets—are driven by operational priorities, as to which management has greater insight than the Board or the Compensation Committee, the Compensation Committee has directed management to interface with the Committee and the compensation consultant to help establish appropriate targets.

Due to the significant cost associated with services provided by a compensation consultant, we may decide not to engage a compensation consultant each year, but rather once every few years. This decision shall be evaluated regularly and will be based on the Compensation Committee's evaluation of whether the prior report obtained, along with increased disclosures of other public companies from our Peer Group relating to executive compensation disclosure, is sufficient to allow them to make informed and reasonable decisions with regard to executive-compensation matters.

Role of our Executives in Setting Compensation

On occasion, the Compensation Committee meets with members of our management team, including Messrs. Kevin Connors and Ron Santilli, to obtain recommendations with respect to Company compensation programs, practices and packages for executives, other employees and directors. Management may make recommendations to the Compensation Committee on all components of compensation. The Compensation Committee considers, but is not bound to and does not always accept, management's recommendations with respect to these matters. The Compensation Committee has the ultimate authority to make decisions with respect to the compensation of our Named Executive Officers and does not delegate any of its compensation functions to others.

Market Benchmarks

In developing its recommendations for annual compensation packages for our Named Executive Officers, our Compensation Committee worked with Mercer to gather market data and identify an appropriate peer group of public companies. The members of that peer group are Athenahealth, Atrion Corporation, Candela (acquired by Syneron Medical, Ltd.), Cryolife, Cynosure, Emageon, Exactech, I-Flow Corporation, Lifecore Biomedical, Medecision, Meridian Bioscience, Osteotech, Palomar Medical Technologies, Quadramed, RTI Biologics, Solta Medical, Transcend Services, and Tutogen Medical (the "Peer Group"). Our Compensation Committee used this data in developing its recommendations for annual compensation for our Named Executive Officers, but also ensured that its recommendations were consistent with the philosophy underlying our compensation programs.

Compensation Components

Our Named Executive Officers are compensated with cash, equity and non-equity incentives, and other customary employee benefits.

Cash Compensation. Cash compensation consists of base salary, participation in a discretionary bonus program and participation in a discretionary profit-sharing plan. Our cash compensation goals for our Named Executive Officers are based upon the following principals:

- Salary should generally be set at or above the 50th percentile of the Peer Group;
- Salary should be positioned to reflect each individual's experience, performance and potential;
- A significant portion of cash compensation should be "at risk;" and
- The amount of discretionary bonuses payable in any quarter is based on revenue growth, compared with the same quarter in the prior year, and the operating profit before stock-based compensation and non-operational expenses, or "Adjusted Operating Profit." Further, discretionary bonuses are payable only if we have an Adjusted Operating Profit for that quarter.

Base Salary and Total Target Cash Compensation. Total target cash compensation for each Named Executive Officer includes his annual base salary, annual target bonus level (described below) and annual profit-sharing payments.

In 2010, our Compensation Committee did not retain a third-party compensation consultant to assist with establishing compensation for our Named Executive Officers. During 2009, our Compensation Committee retained Mercer, to assist it with establishing compensation for our Named Executive Officers and paid them approximately \$68,000 in consulting fees. Mercer does not provide any other services to us other than the services it performs at the request of the Compensation Committee. Mercer assisted the Compensation Committee's executive compensation-setting process by:

- assessing the competitiveness of our compensation arrangements for the Named Executive Officers and making recommendations regarding equity grants to these individuals;
- assessing the competitiveness of our compensation arrangements for the members of our Board and making recommendations regarding the compensation program's design and levels; and
- reviewing and providing comments on the structure of the stock option exchange program.

In 2009, Mercer concluded that our executive total target cash compensation program was well aligned with the practices of the broader medical device industry and direct peers. For 2010, the annual base salary and total target cash compensation of our Named Executive Officers remained the same as it was in 2009.

Discretionary Bonus Program. In addition to base salary compensation, we had a discretionary bonus program in 2010 for our Named Executive Officers and other personnel pursuant to which cash payments may be made quarterly based on the Company's performance in the then-preceding quarter. The annual target bonus levels as a percentage of base salary for the Named Executive Officers effective through December 31, 2010, remained the same as that set by our Board in February 2008 at 60% for Mr. Connors and 45% for Mr. Santilli. Payments made under the discretionary bonus program are at the Board's option. We, however, have historically made payments in accordance with this program.

Target bonuses in 2010 were calculated based upon a matrix of revenue growth and Adjusted Operating Profit. For example, at 10% revenue growth and 10% Adjusted Operating Profit, an individual would receive 100% of his or her target bonus. At 15% revenue growth and 15% Adjusted Operating Profit, an individual would receive 150% of his or her target bonus. There was no bonus earned by each of our Named Executive Officers in 2010 as we did not have a profitable quarter.

Discretionary Profit-Sharing Program. We also have a discretionary profit sharing program for our Named Executive Officers and other employees pursuant to which cash payments may be made quarterly. Target profit-sharing payments are calculated based upon half of the quarterly pre-tax Adjusted Operating Profit percentage (pre-tax Adjusted Operating Profit divided by revenue) multiplied by the Named Executive Officer's gross salary earned during that quarter. There were no profit-sharing payments made to our Named Executive Officers in 2010 because we did not have an Adjusted Operating Profit in any quarter.

Long-Term Incentive Program. We believe that equity-based compensation promotes and encourages long-term successful performance by our Named Executive Officers that is aligned with the organization's goals and the generation of stockholder value. Our equity compensation goals for our Named Executive Officers and others are based upon the following principals:

- Stockholder and executive interests should be aligned;
- Key and high-performing employees, who have a demonstrable impact on our performance and /or stockholder value, should be provided this benefit;
- The program should be structured to provide meaningful retention incentives to participants;
- The equity grants should reflect each individual's experience, performance, potential and be comparable to what the Peer Group grants for the respective position; and
- Actual awards should be tailored to reflect individual performance and attraction/retention goals.

Equity Incentive Compensation. Under our 2004 Equity Incentive Plan, we are permitted to grant stock options, stock appreciation rights, restricted shares, restricted stock units, performance shares, and other stock-based awards. Under that Plan, we grant options to our officers, directors and employees to purchase shares of our common stock at an exercise price equal to the fair market value of such stock on the date of grant. The grant date for stock options to our Named Executive Officers is typically the date of a regularly scheduled board meeting, or, for annual merit grants, on or around June 1 of each year. Our outside directors are granted stock annually on the date of our Annual Meeting of stockholders. We have no program, plan or practice to select option grant dates (or set board meeting and annual stockholder meeting dates) to correspond with the release of material non-public information.

In May 2010, our Compensation Committee, with the approval of our non-employee, outside directors, granted options to Messrs. Connors and Santilli to acquire 120,000 and 55,000 shares of our common stock under our 2004 Equity Incentive Plan, respectively. These equity grants are in the form of stock option awards. At the time these awards were granted, the unvested stock options of each of our Named Executive Officers had exercise prices that were higher than the then-current per-share price of our common stock. Each of the stock option awards had a vesting commencement date of June 1, 2010, a term of seven years, and vests as follows: twelve thirty-sixths of the total number of shares subject to the stock option shall vest one full calendar year following the vesting commencement date of June 1, 2010 and one-thirty-sixth of the total number of shares subject to the stock option shall vest on the last day of each full calendar month thereafter, until all such shares have vested, subject to the option holder continuing to provide services to us through each such date.

Each of the stock option awards described in the preceding paragraph was issued for the purpose of retaining the individual recipient and thus ultimately increasing the value of the Company. In connection with the granting of these stock options, stock options to Messrs. Connors and Santilli to acquire 95,000 and 57,000 shares of our common stock, respectively, that had exercise prices of \$23.75 and \$24.46 — significantly higher than the thencurrent per-share price of our common stock — were cancelled.

In addition, in May 2010, our Compensation Committee, with the approval of our non-employee, outside directors, granted Restricted Stock Units (or RSUs) to Messrs. Connors and Santilli to acquire 33,000 and 22,000 shares of our common stock under our 2004 Equity Incentive Plan, respectively. These RSUs vest as to one-third of the shares on each of June 1, 2010, 2011 and 2012.

We also have a 2004 Employee Stock Purchase Plan that provides eligible employees with the opportunity to purchase shares of our common stock at a 15% discounted price to the lower of the fair market value at either the beginning or the end of the applicable offering period. During 2010, our Chief Financial Officer participated in this Plan, but our Chief Executive Officer did not participate in this Plan.

Benefits. We provide the following benefits to our Named Executive Officers generally on the same basis as the benefits provided to all employees:

- Health, dental and vision insurance;
- Life insurance;
- Short-and long-term disability;
- 401(k) plan, however, in 2009 we discontinued our discretionary employer match on employee 401(k) contributions; and
- Flexible Spending Accounts.

These benefits are consistent with those offered by other companies and specifically with those companies with which we compete for employees.

We have Change of Control and Severance Agreements with each of our Named Executive Officers. The purpose of these agreements is to provide incentives to our Named Executive Officers to continue their employment with the Company and not be distracted by the possibility of loss of employment as a result of an acquisition of the Company or for other reasons.

The Change of Control and Severance Agreements provide that if a Named Executive Officer's employment with the Company is terminated by the Company without Cause or by the executive for Good Reason either prior to 3 months before or after 12 months following a Change of Control (as such capitalized terms are defined in the Change of Control and Severance Agreement) of the Company but not in connection with a Change of Control, the executive will receive, subject to signing a release of claims in favor of the Company, (i) a lump sum severance payment equal to 200% of the annual base salary as in effect immediately prior to such termination for our Chief Executive Officer and 100% of the annual base salary as in effect immediately prior to such termination for our Chief Financial Officer; and (ii) up to 24 months for our Chief Executive Officer and up to 12 months for our Chief Financial Officer of reimbursement for premiums paid for COBRA coverage.

The Change of Control and Severance Agreements also provide that if an executive's employment with the Company is terminated by the Company without Cause or by the executive for Good Reason and such termination occurs within the period beginning 3 months before, and ending 12 months following, a Change of Control of the Company and in connection with a Change of Control, the executive will receive, subject to signing a release of claims in favor of the Company, (i) a lump sum severance payment equal to 200% of the annual base salary as in effect immediately prior to such termination or, if greater, at the level in effect immediately prior to the Change of Control for our Chief Executive Officer and 100% of the annual base salary as in effect immediately prior to such termination or, if greater, at the level in effect immediately prior to the Change of Control for our Chief Financial Officer; (ii) a lump sum severance payment equal to 100% of the executive's annual target bonus for the fiscal year in which the termination occurs or, if greater, executive's annual target bonus in effect immediately prior to the Change of Control; (iii) automatic vesting in full of all outstanding and unvested equity awards held by the executive as of the date of the Change of Control; and (iv) up to 24 months for our Chief Executive Officer and up to 12 months for our Chief Financial Officer of reimbursement for premiums paid for COBRA coverage.

Each Change of Control and Severance Agreement has an initial term of three years, and will extend for an additional year unless the Company or the applicable executive provides written notice at least sixty days prior to the third anniversary of the agreement. For purposes of these agreements, "Cause" shall mean executive's termination only upon (i) executive's willful failure to substantially perform executive's duties (subject to notice and a reasonable period to cure), other than a failure resulting from executive's complete or partial incapacity due to physical or mental illness or impairment; (ii) executive's willful act which constitutes gross misconduct and which is injurious to the Company; (iii) executive's willful breach of a material provision of the Change of Control and Severance Agreement (subject to notice and reasonable period to cure); or (iv) executive's knowing, material and willful violation of a federal or state law or regulation applicable to the business of the Company.

For purposes of these agreements, "Good Reason" shall mean executive's termination of employment within ninety (90) days following the expiration of any cure period following the occurrence of one or more of the following, without executive's consent: (i) a material reduction in executive's authority, duties, or responsibilities relative to duties, position or responsibilities in effect immediately prior to such reduction; (ii) a material reduction in executive's base salary as in effect immediately prior to such reduction; or (iii) a material change in the geographic location at which executive must perform services (in other words, the relocation of executive to a facility that is more than fifty (50) miles from executive's then-current location.

Based on its review of our change-in-control program in 2009, Mercer determined that it is aligned with the practices of our direct peers.

Potential Payments Upon Termination or Change in Control

The following table lists our Named Executive Officers and the estimated amounts they would have become entitled to had their employment with us terminated without Cause or resigns for Good Reason not in connection with a Change of Control on December 31, 2010.

	Estimated Total Value	Estimated Total Value of Health Coverage Continuation	
Name	of Cash Payment		
Kevin P. Connors	\$ 840,000	\$ 15,075	
Ronald J. Santilli	\$ 290,000	\$ 21,242	

The following table lists our Named Executive Officers and the estimated amounts they would have become entitled to had their employment with us terminated without Cause or resigns for Good Reason in connection with a Change of Control on December 31, 2010.

		Estimated	
·	Estimated	Total Value	
	Total Value	of Health	Value of
	of Cash	Coverage	Accelerated
Name	Payment	Continuation	Equity (i)
Kevin P. Connors	\$1,092,000	\$ 15,075	\$ 182,380
Ronald J. Santilli	\$ 420,500	\$ 21,242	\$ 121,587

(1) We estimate the value of acceleration of unvested options and RSUs held by each of our Named Executive Officers based on a share price of \$8.29 per share as of December 31, 2010.

Except for these Change of Control and Severance Agreements, we do not have employment agreements with any of our Named Executive Officers.

Internal Revenue Code Section 162(m) and Limitations on Executive Compensation

Section 162(m) of the United States Internal Revenue Code of 1986, as amended, may limit our ability to deduct for United States federal income tax purposes compensation paid to either our Chief Executive Officer or to other highly paid executive officers in any one fiscal year that is, for each such person, in excess of \$1,000,000. None of our executive officers received any such compensation in excess of this limit during 2010, or any prior year.

Grants of stock options under the 2004 Equity Incentive Plan are not subject to the deduction limitation; however, to preserve our ability to deduct the compensation income associated with options granted to such executive officers pursuant to Section 162(m) of the Internal Revenue Code, our 2004 Equity Incentive Plan provides that no optionee may be granted option(s) to purchase more than 500,000 shares of our common stock in any one fiscal year. However, in the fiscal year in which the optionee is hired, an optionee may be granted an option to purchase up to 1,000,000 shares of our common stock.

Securities Authorized for Issuance Under Equity Compensation Plans

Our stockholders approved each of our equity compensation plans, including a 2008 amendment to our 2004 Equity Incentive Plan. The following table provides information regarding common stock that may be issued upon the exercise of options and restricted stock units under our 2004 Equity Incentive Plan as of December 31, 2010.

Plan category	Number of securities to be issued upon exercise of outstanding options, warrants and rights (a)	Weighted-average exercise price of outstanding options, warrants and rights (b)		remaining available for future issuance under equity compensation plans (excluding securities reflected in column (a)) (c)	
Equity compensation plans approved by security holders	3,296,419	\$	10.93	1,005,447	
by security holders	3,296,419	\$	10.93	1,005,447	

Number of securities

Summary Compensation Table

The following table sets forth summary compensation information for the years ended December 31, 2010, 2009 and 2008 for our Chief Executive Officer and Chief Financial Officer. We refer to these persons as our Named Executive Officers elsewhere in this proxy statement. Except as provided below, none of our Named Executive Officers received any other compensation required to be disclosed by law or in excess of \$10,000 annually.

Name and Principal Position Kevin P. Connors President and Chief	Salary Bonus ⁽¹⁾ Awards ⁽²⁾ Co		Non-Equity Incentive Plan Compensation ⁽³⁾	All Other Compensation (4)	Total	
Executive Officer 2010	\$ 420,000 420,000 420,000	\$ — 18,454 53,086	\$ 729,772 481,284 708,145	\$ <u> </u>	\$	\$ 1,149,772 919,738 1,198,900
Ronald J. Santilli Executive Vice President and Chief Financial Officer 2010	\$ 290,000	\$ —	\$ 396,546	\$ —	\$ —	\$ 686,546
2009 2008	290,000 290,000	10,012 28,513	220,589 339,765		10,350	520,601 668,628

⁽¹⁾ Amounts represent a discretionary bonus and profit sharing earned for each of the years in 2010, 2009 and 2008.

Grants of Plan-Based Awards

The following table lists grants of plan-based stock options and restricted stock unit awards made to our Named Executive Officers in 2010 at their related grant date fair value calculated in accordance with ASC Topic 718.

⁽²⁾ Amounts shown in this column are the aggregate grant date fair value of stock awards granted during each of the years in 2010, 2009 and 2008 calculated in accordance with ASC Topic 718. See Note 5 of the Consolidated Notes to Financial Statements included in our Annual Report on Form 10-K for the year ended December 31, 2010 filed with the SEC on March 15, 2011 for a discussion of valuation assumptions for stock-based compensation.

Amounts represent non-cash benefit associated with a company sponsored, non-business, event for achieving sales targets in accordance with our commission incentive plan.

⁽⁴⁾ Amount represents 401(k) employer-match contributions and service award, where applicable. In 2009, we discontinued our discretionary employer match on employee 401(k) contributions.

			Estimated Future Payouts Under Non-Equity Incentive Plan Awards			All Other Option Awards: Number of	Exercise or Base Price	Grant Date Fair Value of
Name	Equity Instrument	Grant Date	Threshold	Target	Maximum	Securities Underlying Options	of Option Awards (2)	Stock Option Awards (1)
Kevin P. Connors President and Chief								
Executive Officer			_	_	_			
	Options	05/14/2010				120,000	\$ 10.24	\$ 445,452
	Options	05/14/2010				(95,000)	(24.05)	(53,600)
· ·	RSUs	05/14/2010				33,000	10.24	337,920
	Total					58,000		729,772
Ronald J. Santilli Chief Financial Officer					* .		•	
and Executive Vice President		05/14/2010	_		_			
	Options	05/14/2010				55,000	\$ 10.24	\$ 204,166
	Options	05/14/2010			*. * *	(57,000)	(24.02)	(32,900)
	RSUs	05/14/2010				22,000	10.24	225,280
	Total					20,000	* * *	396,546

⁽¹⁾ Amounts reflect grant date fair value of equity awards calculated in accordance with ASC Topic 718. See Note 5 of the Notes to Consolidated Financial Statements included in our Annual Report on Form 10-K for the year ended December 31, 2010 filed with the SEC on March 15, 2011 for a discussion of valuation assumptions for stock-based compensation.

Equity Incentive Awards Outstanding

The following table lists the outstanding equity incentive awards held by our Named Executive Officers as of December 31, 2010.

		Stock Awards (4)							
Name	Number of Securities Underlying Unexercised Earned Options	Number of Securities Underlying Unexercised Unearned Options	ecurities Iderlying exercised Option nearned Exercise		Option Expiration Date	Number of Shares or Units of Stock that Have Not Vested	Market Value of Shares or Units of Stock that Have Not Vested		Date Awards Will be Fully Vested
Kevin P. Connors	40,000		\$	2.50	6/08/2011				
President and Chief Executive		· .							
Officer	3,333	· —		4.25	8/13/2013				
	30,000	· —		20.25	7/28/2015				
	20,813	12,487(1)		10.43	5/28/2015				
•		$100,000^{(2)}$		10.43	5/28/2015				
	60,000	$60,000^{(3)}$		8.66	6/08/2016				
	· —	$120,000^{(3)}$		10.24	5/14/2017				
						22,000	\$	182,380	6/01/2012
Ronald J. Santilli	20,000	_	\$	5.50	9/24/2011	- -			
Executive	3,372	_		4.25	8/07/2012				
Vice President	14,753	_		4.25	8/13/2013				
	10,000	_		13.30	7/20/2014				
	15,000	_		20.25	7/28/2015				
	8,563	5,137(1)		10.43	5/28/2015				
	_	50,000(2)		10.43	5/28/2015				
	27,501	27,499 ⁽³⁾		8.66	6/08/2016				
		55,000 ⁽³⁾		10.24	5/14/2017				
		,-				14,666	\$	121,581	6/01/2012

⁽¹⁾ One-quarter (1/4th) of the shares underlying each of these options vest on the one year anniversary of the vesting commencement date and 1/48th of the underlying shares vest each month thereafter.

⁽²⁾ The per-share prices were the closing price of our common stock on the respective dates of grant.

^{100%} of the shares underlying each of these options vest on the three year anniversary of the vesting commencement date.

⁽²⁾ (3) One-third (1/3rd) of the shares underlying each of these options vest on the one year anniversary of the vesting commencement date and 1/36th of the underlying shares vest each month thereafter.

⁽⁴⁾ One-half of the shares underlying each of these awards will vest on June 1, 2011 and 2012.

Options Exercised and Stock Vested

The following table lists the options exercised by, and stock vested to, our Named Executive Officers in the year ended December 31, 2010.

	Option	Awards	Stock Awards			
Name	Number of Shares Acquired on Exercise	Value Realized on Exercise (1)	Number of Shares Acquired on Vesting	Value Realized Upon Vesting (2)		
Kevin P. Connors President and Chief Executive Officer Ronald J. Santilli Chief Financial Officer and Executive	50,000	\$ 347,500	11,000			
Vice President	- -		7,334	\$ 64,613		

⁽¹⁾ Represents the excess of fair market value of the shares exercised on the exercise date over the aggregate exercise price for such shares.

⁽²⁾ These shares were originally issued by us pursuant to RSUs granted in May 2010. On each vesting date, the RSU has a value equal to the fair market value of our common stock on the date of vesting.

COMPENSATION COMMITTEE REPORT (1)

The Compensation Committee has reviewed and discussed the Compensation Discussion and Analysis required by Item 402(b) of SEC Regulation S-K with management. Based on such review and discussions, the Compensation Committee recommended to the Board of Directors that the Compensation Discussion and Analysis be included in Cutera's proxy statement.

The foregoing report is provided by the undersigned members of the Compensation Committee.

David B. Apfelberg Timothy J. O'Shea Jerry P. Widman

⁽¹⁾ The material in this report is not deemed soliciting material or filed with the SEC and is not to be incorporated by reference in any filing of the Company under the Securities Act of 1933, as amended, or the Securities Exchange Act of 1934, as amended, whether made before or after the date of this Proxy Statement and irrespective of any general incorporation language in those filings.

OTHER MATTERS

We are not aware of any other business to be presented at the meeting. As of the date of this proxy statement, no stockholder had advised us of the intent to present any business at the meeting. Accordingly, the only business that our Board of Directors intends to present at the meeting is as set forth in this proxy statement.

If any other matter or matters are properly brought before the meeting, the proxies will use their discretion to vote on such matters in accordance with their best judgment.

By order of the Board of Directors,

Kevin P. Connors

President and Chief Executive Officer

Brisbane, California April 26, 2011

THIS PROXY IS SOLICITED ON BEHALF OF THE BOARD OF DIRECTORS OF CUTERA, INC. 2011 ANNUAL MEETING OF STOCKHOLDERS

The undersigned stockholder of Cutera, Inc., a Delaware corporation, hereby acknowledges receipt of the Notice of Annual Meeting of Stockholders and Proxy Statement each dated April 26, 2011 and hereby appoints David B. Apfelberg (the lead independent director on our Board) and Ronald J. Santilli (our Chief Financial Officer), each as proxy and attorney-in-fact, with full power of substitution, on behalf and in the name of the undersigned to represent the undersigned at the 2011 Annual Meeting of Stockholders of Cutera, Inc. to be held on June 14, 2011 at 10:00 a.m., local time, at Cutera's offices located at 3240 Bayshore Blvd., Brisbane, California 94005-1021, and at any postponement or adjournment thereof, and to vote all shares of common stock which the undersigned would be entitled to vote if then and there personally present, on the matters set forth below:

SEE REVERSE SIDE

FOLD AND DETACH HERE

. 1		mark your votes as	_	_		
1.Election of FOR WITHHOLD 2. Directors	A non-binding advisory vote on the approval of executive compensation.	indicated FOR	AGAINST	ABSTAIN		
	2.3 (2.1)	· 🗖	_			
CLASS I 3. NOMINEES:	A non-binding advisory vote on the frequency of executive compensation voting.	Year 2 Year	3 Year	ABSTAIN		
Kevin P. Connors	compensation voting.	•				
David A. Gollnick						
THE 4. STOCKHOLDER MAY WITHHOLD AUTHORITY TO VOTE FOR ANY NOMINEE BY STRIKING OUT THE INDIVIDUAL'S NAME ABOVE	Ratify the appointment of PricewaterhouseCoopers LLP as the Independent Registered Public Accounting Firm of the Company for the fiscal year ending December 31, 2011.	FOR	AGAINST	ABSTAIN		
TWEND THE VE	•					
THIS PROXY WILL BE VOTED AS DIRECTED OR, IF NO CONTRARY DIRECTION IS INDICATED, WILL BE VOTED AS FOLLOWS: (1) FOR THE ELECTION OF THE NOMINATED CLASS I DIRECTORS; (2) FOR THE APPROVAL, BY NON-BINDING VOTE, OF EXECUTIVE COMPENSATION; (3) EVERY 3 YEARS FOR THE APPROVAL, BY NON-BINDING VOTE, OF TRIENNIAL EXECUTIVE COMPENSATION VOTING; (4) FOR THE RATIFICATION OF THE APPOINTMENT OF PRICEWATERHOUSECOOPERS LLP AS OUR INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM; AND (5) AS THE PROXY HOLDERS DEEM ADVISABLE ON SUCH OTHER MATTERS AS MAY COME BEFORE THE MEETING.						
PLEASE SIGN EXACTLY AS YOUR NAME APPEARS HEREON. IF THE STOCK IS REGISTERED IN THE NAME OF TWO OR MORE PERSONS, EACH SHOULD SIGN. EXECUTORS, ADMINISTRATORS, TRUSTEES, GUARDIANS AND ATTORNEYS-IN-FACT SHOULD ADD THEIR TITLES. IF SIGNER IS A CORPORATION, PLEASE GIVE FULL CORPORATE NAME AND HAVE A DULY AUTHORIZED OFFICER SIGN, STATING TITLE. IF SIGNER IS A PARTNERSHIP, PLEASE SIGN IN PARTNERSHIP NAME BY AUTHORIZED PERSON.						
PLEASE SIGN, DATE AND PROMPTLY RETURN THIS PROXY IN THE ENCLOSED RETURN ENVELOPE, WHICH IS POSTAGE PREPAID IF MAILED IN THE UNITED STATES.						
SIGNATURE(S)	SIGNATURE(S)	D	ATE:			
NOTE: This Proxy should be marked, signeturned promptly in the enclosed envelo	gned by the stockholder(s) exactly	as his or her na	me appears	hereon, and		

Please

held by joint tenants or as community property, both should sign.

UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

FORM 10-K

ANNUAL REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

For fiscal year ended December 31, 2010

Commission file number: 000-50644



Cutera, Inc. (Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation or organization)

77-0492262 (I.R.S. Employer Identification Number)

3240 Bayshore Blvd. Brisbane, California 94005 (415) 657-5500

(Address, including zip code, and telephone number, including area code, of registrant's principal executive offices)

Securities registered pursuant to Section 12(b) of the Act:

Title of Each Class

Name of Each Exchange on Which Registered

Common Stock, \$0.001 par value per share

The NASDAQ Stock Market, LLC

Securities Registered Pursuant to Section 12(g) of the Act: None

Indicate by check mark if the registrant is a well-known seasoned issuer, as defined in Rule 405 of the Securities Act. Yes 🗆 No 🗵

Indicate by check mark if the registrant is not required to file reports pursuant to Section 13 or Section 15(d) of the Act. Yes 🗆 No 🗵

Indicate by check mark whether the registrant (1) has filed all reports required to be filed by Section 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period than the registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days. Yes \boxtimes No \square

Indicate by check mark whether the registrant has submitted electronically and posted on its corporate Web site, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T (§232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit and post such files). Yes \square No \square

Indicate by check mark if disclosure of delinquent filers pursuant to Item 405 of Regulation S-K is not contained herein, and will not be contained, to the best of the registrant's knowledge, in definitive proxy or information statements incorporated by reference in Part III of this Form 10-K or any amendment to this Form 10-K.

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, or a non-accelerated filer. See definition of "accelerated filer and large accelerated filer" in Rule 12b-2 of the Exchange Act (check one):

reporting company)

Large accelerated filer [7]

Accelerated

filer 🖾

Non-accelerated filer (Do not check if a smaller

Smaller reporting company □

Indicate by check mark whether registrant is a shell company (as defined in Rule 12b-2 of the Exchange Act). Yes 🗆 No 🗵

The aggregate market value of the registrant's common stock, held by non-affiliates of the registrant as of June 30, 2010 (which is the last business day of registrant's most recently completed second fiscal quarter) based upon the closing price of such stock on the NASDAQ Global Select Market on that date, was approximately \$72 million. For purposes of this disclosure, shares of common stock held by entities and individuals who own 5% or more of the outstanding common stock and shares of common stock held by each officer and director have been excluded in that such persons may be deemed to be "affiliates" as that term is defined under the Rules and Regulations of the Securities Exchange Act of 1934. This determination of affiliate status is not necessarily conclusive.

The number of shares of Registrant's common stock issued and outstanding as of February 28, 2011 was 13,669,258.

DOCUMENTS INCORPORATED BY REFERENCE

Part III incorporates by reference certain information from the registrant's definitive proxy statement for the 2011 Annual Meeting of Stockholders.

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ITEM 1. BUSINESS

We are a global medical device company headquartered in Brisbane, California specializing in the design, development, manufacture, marketing and servicing of laser and light-based aesthetics systems for practitioners worldwide. We offer easy-to-use products based on five platforms—CoolGlide[®], Xeo[®], Solera[®], GenesisPlusTM and Excel VTM — each of which enable physicians and other qualified practitioners to perform safe and effective aesthetic procedures for their customers.

- CoolGlide- In March 2003, our first product platform, CoolGlide, was launched. This platform offers
 laser applications for hair removal, treatment of a range of vascular lesions, including leg and facial veins,
 and Laser Genesis—a skin rejuvenation procedure that reduces fine lines, reduces pore size and improves
 skin texture.
- **Xeo-** In 2003, we introduced the Xeo platform, which can combine pulsed light and laser applications in a single system. The Xeo is a fully upgradeable platform on which a customer can use every application that we offer to remove unwanted hair, treat vascular lesions and rejuvenate the skin by treating discoloration, improving texture, reducing pore size and treating fine lines and laxity. This product platform represents the largest contributor to our Product and Upgrade revenue.
- Solera- In 2004, we introduced the Solera platform, a compact tabletop system designed to support a single technology platform. Solera systems use either infrared (Solera Titan) or pulsed light (Solera Opus) and can be used to remove unwanted hair, treat vascular lesions and rejuvenate the skin. The Solera Opus can support one or more pulsed light applications in a single system.
- GenesisPlus- In 2010, we introduced the GenesisPlus platform, which is a dedicated laser based system for performing skin rejuvenation procedures and toe nail fungus removal (CE mark approved and FDA clearance pending). This system has a hand piece that includes real time temperature monitoring of the treatment area, as well as a non-contact distance gauge using two aiming beams, for improving the clinical result of the treatment.
- Excel V- In February 2011, we introduced our Excel V platform, a high-performance, vascular platform designed specifically for the core-market of Dermatologists and Plastic Surgeons. This platform provides a combination of the 532 nm green laser with Cutera's award winning 1064 nm Nd:YAG technology, to provide a single, compact and efficient system that treats the entire range of cosmetic vascular conditions, without the need for costly consumables.

Each of our laser and light-based platforms consists of one or more hand pieces and a console that incorporates a universal graphic user interface, a laser or light-based module, control system software and high voltage electronics. However, depending on the application, the laser or light-based module is sometimes instead contained in the hand piece. A description of each of our hand pieces, and the aesthetic conditions they are designed to treat, are contained in the section entitled "Products," below.

We offer our customers the ability to select the systems and applications that best fit their practice and to subsequently upgrade their systems to add new applications. This upgrade path allows our customers to cost-effectively build their aesthetic practices and provides us with a source of recurring revenue.

In addition to systems and upgrades, we generate revenue from the sale of post warranty services, Titan hand piece refills, and Dermal filler and cosmeceuticals.

The Structure of Skin and Conditions that Affect Appearance

The skin is the body's largest organ and is comprised of layers called the epidermis and dermis. The epidermis is the outer layer, and serves as a protective barrier for the body. It contains cells that determine pigmentation, or skin color. The underlying layer of skin, the dermis, contains hair follicles and large and small blood vessels that are found at various depths below the epidermis. Collagen, also found within the dermis, provides strength and flexibility to the skin.

Many factors, such as age, smoking and sun damage, can result in aesthetically unpleasant changes in the appearance of the skin. These changes can include:

- Undesirable hair growth;
- Enlargement or swelling of blood vessels due to circulatory changes that become visible at the skin's surface in the form of unsightly veins;
- Deterioration of collagen, which weakens the skin, leading to uneven texture, increased pore size, wrinkles and laxity; and
- Uneven pigmentation or sun spots due to long-term sun exposure.

People with unwanted hair or any of the above-mentioned skin conditions often seek aesthetic treatments to improve their appearance.

The Market for Non-Surgical Aesthetic Procedures

The market for non-surgical aesthetic procedures has grown significantly over the past several years. The American Society of Plastic Surgeons estimates that in 2010 there were over 11.5 million minimally-invasive aesthetic procedures performed, a 5% increase over 2009 and a 110% increase over 2000. We believe there are several factors contributing to the growth of these aesthetic procedures, including:

- Aging of the U.S. Population- The "baby boomer" demographic segment ages 46 to 64 in 2010 represented approximately 80 million people, or 26%, of the U.S. population in 2010. The size of this aging segment, and its desire to retain a youthful appearance, has contributed to the growth for aesthetic procedures.
- Broader Range of Safe and Effective Treatments- Technical developments have led to safe, effective, easy-to-use and low-cost treatments with fewer side effects, resulting in broader adoption of aesthetic procedures by practitioners. In addition, technical developments have enabled practitioners to offer a broader range of treatments. These technical developments have reduced the required treatment and recovery times, which in turn have led to greater patient demand.
- Broader Base of Customers- Managed care and government payer reimbursement restrictions in the United States, and similar payment related constraints outside the United States, may help motivate qualified practitioners from differing specialties to establish or expand their elective aesthetic practices with procedures that are paid for directly by patients. As a result, in addition to the core users such as dermatologists and plastic surgeons, many other non-core practitioners, such as gynecologists, family practitioners, primary care physicians, physicians offering aesthetic treatments in non-medical offices, and other qualified practitioners are offering aesthetic procedures.

Non-Surgical Aesthetic Procedures for Improving the Skin's Appearance and Their Limitations

Many alternative therapies are available for improving a person's appearance by treating specific structures within the skin. These procedures utilize injections or abrasive agents to reach different depths of the dermis and the epidermis. In addition, non-invasive and minimally-invasive treatments have been developed that employ laser and light-based technologies to achieve similar therapeutic results. Some of these more common therapies and their limitations are described below.

Hair Removal- Techniques for hair removal include waxing, depilatories, tweezing, shaving, electrolysis and laser and light-based hair removal. The only techniques that provide a long-lasting solution are electrolysis and light-based hair removal. Electrolysis is usually painful, time-consuming and expensive for large areas, but is the most common method for removing light-colored hair. During electrolysis, an electrologist inserts a needle directly into a hair follicle and activates an electric current in the needle. Since electrolysis only treats one hair follicle at a time, the treatment of an area as small as an upper lip may require numerous visits and many hours of treatment. In addition, electrolysis can cause blemishes and infection related to needle use.

Leg and Facial Veins- The current aesthetic treatment methods for leg and facial veins include sclerotherapy and laser and light-based treatments. With these treatments, patients seek to eliminate visible veins and improve overall skin appearance. Sclerotherapy requires a skilled practitioner to inject a saline or detergent-based solution into the target vein, which breaks down the vessel causing it to collapse and be absorbed into the body. The need to correctly position the needle on the inside of the vein makes it difficult to treat smaller veins, which limits the treatment of facial vessels and small leg veins. The American Society of Plastic Surgeons estimates that approximately 365,000 sclerotherapy procedures were performed in 2010.

Skin Rejuvenation- Skin rejuvenation treatments include a broad range of popular alternatives, including Botox and collagen injections, chemical peels, microdermabrasions, radiofrequency treatments and lasers and light-based treatments. With these treatments, patients hope to improve overall skin tone and texture, reduce pore size, tighten skin and remove other signs of aging, including mottled pigmentation, diffuse redness and wrinkles. All of these procedures are temporary solutions and must be repeated within several weeks or months to sustain their effect, thereby increasing the cost and inconvenience to patients. For example, the body absorbs Botox and collagen and patients require supplemental injections every three to six months to maintain the benefits of these treatments.

Some skin rejuvenation treatments, such as chemical peels and microdermabrasions, can have undesirable side effects. Chemical peels use acidic or caustic solutions to peel away the epidermis, and microdermabrasion generally utilizes sand crystals to resurface the skin. These techniques can lead to stinging, redness, irritation and scabbing. In addition, more serious complications, such as changes in skin color, can result from deeper chemical peels. Patients that undergo these deep chemical peels are also advised to avoid exposure to the sun for several months following the procedure. The American Society of Plastic Surgeons estimates that in 2010, approximately 5.4 million injections of Botox and 1.8 million injections of collagen and other soft-tissue fillers were administered; and 1.1 million chemical peels and 825,000 microdermabrasion procedures were performed.

In radiofrequency tissue tightening, energy is applied to heat the dermis of the skin with the goal of shrinking and tightening the collagen fibers. This approach may result in a more subtle and incremental change to the skin than a surgical facelift. Drawbacks to this approach may include surface irregularities that may however resolve over time, and the risk of burning the treatment area.

Laser and light-based non-surgical treatments for hair removal, veins and skin rejuvenation are discussed in the following section and in the section entitled "Our Applications and Procedures," below.

Laser and Light-Based Aesthetic Treatments

Laser and light-based aesthetic treatments can achieve therapeutic results by affecting structures within the skin. The development of safe and effective aesthetic treatments has created a well-established market for these procedures.

Ablative skin resurfacing is a method of improving the appearance of the skin by removing the outer layers of the skin. Ablative skin resurfacing procedures are considered invasive or minimally invasive, depending on how much of the epidermis is removed during a treatment. Non-ablative skin resurfacing is a method of improving the appearance of the skin by treating the underlying structure of the skin without damaging the outer layers of the skin. Practitioners can use laser and light-based technologies to selectively target hair follicles, veins or collagen in the dermis, as well as cells responsible for pigmentation in the epidermis, without damaging surrounding tissue. They can also use these technologies to safely remove portions of the epidermis and deliver heat to the dermis as a means of generating new collagen growth.

Safe and effective laser and light-based treatments require an appropriate combination of the following four parameters:

- Energy Level- the amount of light emitted to heat a target;
- Pulse Duration- the time interval over which the energy is delivered;
- Spot Size- the diameter of the energy beam, which affects treatment depth and area; and
- Wavelength- the color of light, which impacts the effective depth and absorption of the energy delivered.

For example, in the case of hair removal, by utilizing the correct combination of these parameters, a practitioner can use a laser or other light source to selectively target melanin within the hair follicle to absorb the laser energy and destroy the follicle, without damaging other delicate structures in the surrounding tissue. Wavelength and spot size permit the practitioner to target melanin in the base of the hair follicle, which is found in the dermis. The combination of pulse duration and energy level may vary, depending upon the thickness of the targeted hair follicle. A shorter pulse length with a high energy level is optimal to destroy fine hair, whereas coarse hair is best treated with a longer pulse length with lower energy levels. If treatment parameters are improperly set, non-targeted structures within the skin may absorb the energy thereby eliminating or reducing the therapeutic effect. In addition, improper setting of the treatment parameters or failure to protect the surface of the skin may cause burns, which can result in blistering, scabbing and skin discoloration.

Technology and Design of Our Systems

Our unique CoolGlide, Xeo, Solera, GenesisPlus and Excel V platforms provide the long-lasting benefits of laser and light-based aesthetic treatments. Our technology allows for a combination of a wide variety of applications available in a single system. Key features of our solutions include:

- Multiple Applications Available in a Single System- Our systems comprise of multi-applications that
 enable practitioners to perform multiple aesthetic procedures using a single device. These procedures
 include hair removal, vascular treatments and skin rejuvenation including the treatment of
 discoloration, laxity, fine lines, pore size and uneven texture. Because practitioners can use our systems
 for multiple indications, the cost of a unit may be spread across a potentially greater number of patients
 and procedures and therefore may be more rapidly recovered.
- Technology and Design Leadership- We offer innovative laser and light-based solutions for the aesthetic market. Our laser technology combines long wavelength, adjustable energy levels, variable spot sizes and a wide range of pulse durations, allowing practitioners to customize treatments for each patient and condition. Our proprietary pulsed light hand pieces for the treatment of discoloration, hair removal and vascular treatments optimize the wavelength used for treatments and incorporate a monitoring system to increase safety. Our Titan hand pieces utilize a novel light source that had not been previously used for aesthetic treatments. Our Pearl and Pearl Fractional hand pieces, with proprietary YSGG technology, represent the first application of the 2790 nm wavelength for minimally-invasive cosmetic dermatology. Further, our GenesisPlus platform for performing skin rejuvenation procedures and toenail fungus removal (CE mark approved and FDA clearance pending) has a hand piece that includes real time temperature monitoring of the treatment area, as well as a non-contact distance gauge using two aiming beams, for improving the clinical result of the treatment.
- Upgradeable Platform- We have designed some of our products to allow our customers to costeffectively upgrade to our multi-application systems (Solera and Xeo), which provide our customers with
 the option to add additional applications to their existing systems and provides us with a source of
 recurring revenue. We believe that product upgradeability allows our customers to take advantage of our
 latest product offerings and provide additional treatment options to their patients, thereby expanding the
 opportunities for their aesthetic practices.
- Treatments for Broad Range of Skin Types and Conditions- Our products remove hair safely and effectively on patients of all skin types, including harder-to-treat patients with dark or tanned skin. In addition, the wide parameter range of our systems allows practitioners to effectively treat patients with both fine and coarse hair. Practitioners may use our products to treat spider and reticular veins (unsightly small veins in the leg) and small facial veins; perform skin rejuvenation procedures for discoloration, texture, pore size, fine lines, and laxity on any type of skin; and treat toe nail fungus (CE mark approved and FDA clearance pending). The ability to customize treatment parameters enables practitioners to offer safe and effective therapies to a broad base of their patients.

• Ease of Use- We design our products to be easy to use. Our proprietary hand pieces are lightweight and ergonomic, minimizing user fatigue, and allow for clear views of the treatment area, reducing the possibility of unintended damage and increasing the speed of application. Our control console contains a universal graphic user interface with three simple, independently adjustable controls from which to select a wide range of treatment parameters to suit each patient's profile. The clinical navigation user interface on the Xeo platform provides recommended clinical treatment parameter ranges based on patient criteria entered. And our Pearl and Pearl Fractional hand pieces include a scanner with multiple scan patterns to allow simple and fast treatments of the face. Risks involved in the use of our products include risks common to other laser and light-based aesthetic procedures, including the risk of burns, blistering and skin discoloration.

Strategy

Our goal is to maintain and expand our position as a leading, worldwide, provider of energy-based aesthetic devices and complementary aesthetic products by executing the following strategies:

- Continue to Expand our Product Offering- Though we believe that our current portfolio of products is comprehensive, our research and development group has a pipeline of potential products under development that we expect to commercialize in the future. In addition to products in the laser and light-based aesthetic market, we are expanding our product offering into other complementary aesthetic applications, such as dermal fillers and cosmeceuticals. Such products will allow us to leverage our existing customer call points, and provide us with new customer call points, to generate additional revenue, which will enhance the productivity of our distribution channels.
- Increasing Revenue and Improving Productivity- We believe that the market for aesthetic systems will continue to offer growth opportunities in the future, even though our revenue declined by 1% in 2010, compared with 2009. We continue to build brand-recognition, add additional products to our international distribution channel and remain focused on enhancing our global distribution network, all of which we expect will increase our revenue. In addition, we plan to grow our U.S. revenue by leveraging our relationship with PSS World Medical Shared Services, Inc., or PSS—a wholly-owned subsidiary of PSS World Medical—that operates medical supply distribution service centers with over 700 sales consultants serving physician offices throughout the United States. In 2010, we expanded our senior sales management to provide increased focus to our distribution channels and improve sales productivity.
- Increasing Focus on Practitioners with Established Medical Offices— We believe there is growth opportunity in targeting our products to a broad customer base. However, in response to the recent global recession, we shifted our focus to the core practitioners and physicians with established medical offices. We believe that our customer success is largely dependent upon having an existing medical practice, in which our systems provide incremental revenue sources to augment their practice revenue.
- Leveraging our Installed Base with Sales of Upgrades- In February 2011, we introduced the Excel V and in 2010, we introduced GenesisPlus both stand alone platforms. However in the past, we have introduced new products that allowed existing customers to upgrade their previously purchased systems to offer additional capabilities. We believe that providing upgrades to our existing installed base of customers continues to represent a potentially significant opportunity for recurring revenue. We also believe that our upgrade program aligns our interest in generating revenue with our customers' interest in improving the return on their investment by expanding the range of applications that can be performed with their existing systems. In 2011, we plan on continuing to market upgrades to our installed base of customers with applications such as Pearl and Pearl Fractional, Titan and other flash lamp hand pieces.
- Generating Revenue from Services and Refillable Hand Pieces- Our Titan hand pieces and pulsed-light
 hand pieces are refillable products, which provide us with a source of recurring revenue from our existing
 customers. We offer post-warranty services to our customers either through extended service contracts to
 cover preventive maintenance or through direct billing for parts and labor. These post-warranty services
 serve as additional sources of recurring revenue.

Products

Our CoolGlide, Xeo, Solera, GenesisPlus and Excel V platforms allow for the delivery of multiple laser and light-based aesthetic applications from a single system. With our Xeo and Solera platforms, practitioners can purchase customized systems with a variety of our multi-technology applications.

The following table lists our products and each checked box represents the incremental applications that were added to the respective platforms in the years noted.

Applications:				Hair Removal:	Vascular Lesions:	Skin Rejuvenation		
System Platforms:	Products:	Year:	Energy Source:			Dyschromia:	Texture, Lines and Wrinkles:	Skin Laxity:
CoolGlide	CV	2000	a	X				
	Excel	2001	a		x			
	Vantage	2002	a				x	
Xeo:	Nd:YAG	2003	a	X	x		x	
	OPS600	2003	b			X		
	LP560	2004	b			X		
	Titan S	2004	С					X
	ProWave 770	2005	b	X				
	AcuTip 500	2005	b		x			
	Titan V/XL	2006	c		,			X
	LimeLight	2006	ь			X		
	Pearl	2007	d·	<i>:</i> ,		X	X	
	Pearl Fractional	2008	d ·	•			X	
Solera	Titan S	2004	c · ·					x
	ProWave 770	2005	b	X				
	OPS 600	2005	ъ			X		
	LP560	2005	b			X		
	AcuTip 500	2005	b		x			
	Titan V/XL	2006	c					x
	LimeLight	2006	b			X		
Genesis Plus		2010	a				X	
Excel V		2011	e		X	x	X	

Energy Source: a. 1064nm Nd:YAG laser; b. flashlamp; c. Infrared laser; d. 2790 nm YSGG laser; e. combined frequency 532 nm and 1064 nm Nd:YAG laser

Each of our products consists of a control console and one or more hand pieces, depending on the model.

Control Console

Our control console includes a universal graphic user interface, control system software and high voltage electronics. All CoolGlide systems, GenesisPlus, Excel V and some models of the Xeo platform, include our laser module which consists of electronics, a visible aiming beam, a focusing lens, and an Nd:YAG and/or flashlamp laser that functions at wavelengths that permit penetration over a wide range of depths and is effective across all skin types. The interface allows the practitioner to set the appropriate laser or flashlamp parameters for each procedure through a user-friendly format. The control system software ensures that the operator's instructions are properly communicated from the graphic user interface to the other components within the system. Our high voltage electronics produce over 10,000 watts of peak laser energy, which permits therapeutic effects at short pulse durations. Our Solera console platform comes in two configurations—Opus and Titan—both of which include a universal graphic user interface, control system software and high voltage electronics. The Solera Opus console is designed specifically to drive our flashlamp hand pieces while the Solera Titan console is designed specifically to drive the Titan hand pieces. The control system software is designed to ensure that the operator's instructions are properly communicated from the graphical user interface to the other components within the system and includes real-time calibration to control the output energy as the pulse is delivered during the treatment.

Hand Pieces

1064 nm Nd:YAG Hand Piece- Our 1064nm Nd:YAG hand piece delivers laser energy to the treatment area for hair removal, leg and facial vein treatment, and skin rejuvenation procedures to treat skin texture and fine lines, and reduce pore size. The 1064nm Nd:YAG hand piece consists of an energy-delivery component, consisting of an optical fiber and lens, and a copper cooling plate with imbedded temperature monitoring. The hand piece weighs approximately 14 ounces, which is light enough to be held with one hand. The lightweight nature and ergonomic design of the hand piece allows the operation of the device without user fatigue. Its design allows the practitioner an unobstructed view of the treatment area, which reduces the possibility of unintended damage to the skin and can increase the speed of treatment. The 1064nm Nd:YAG hand piece also incorporates our cooling system, providing integrated pre- and post cooling of the treatment area through a temperature-controlled copper plate to protect the outer layer of the skin. The hand piece is available in either a fixed 10 millimeter spot size for our CoolGlide CV system, or a user-controlled variable 3, 5, 7 or 10 millimeter spot size for our CoolGlide Excel and CoolGlide Vantage systems.

Excel V Hand Piece- The Excel V system introduced in February 2011 delivers 1064 nm and 532 nm laser energy to the treatment area for vascular treatments. The Excel V system includes two hand pieces, both consisting of an energy-delivery component, consisting of an optical fiber and lens. One hand piece includes a sapphire window cooling plate with temperature monitoring. The second hand piece does not have a cooling plate and includes a non-contact temperature sensor to monitor the treatment area temperature. In addition, this second hand piece includes two aiming beams that facilitate consistent treatments by maintaining the correct distance of the hand piece to the skin. Both hand pieces offer a spot size range from 1.5 to 12 mm in 0.1 mm increments. Each hand piece is capable of delivering either the 1064 nm or 532 nm laser energy.

GenesisPlus Hand Piece- Our GenesisPlus system launched in 2010 delivers 1064 nm laser energy to the treatment area for skin rejuvenation procedures to treat skin texture and fine lines, and reduce pore size. This 1064nm Nd:YAG hand piece consists of an energy-delivery component, consisting of an optical fiber and lens but is lighter since it does not include a copper cooling plate. The hand piece does include a non-contact temperature sensor to monitor the treatment area temperature. In addition, the hand piece includes two aiming beams that facilitate consistent treatments by maintaining the correct distance of the hand piece to the skin. This hand piece offers a single 5 mm spot size.

Pulsed Light Hand Piece- The LP560, ProWave 770, AcuTip 500 and LimeLight hand pieces are designed to produce a pulse of light over a wavelength spectrum to treat discoloration, including pigmented lesions, such as age and sun spots, hair removal and superficial facial vessels. The hand pieces each consist of a custom flashlamp, proprietary wavelength filter, closed-loop power control and embedded temperature monitor, and weigh approximately 13 ounces. The filter in the AcuTip 500 eliminates long and short wavelengths, transmitting only the therapeutic range required for safe and effective treatment. The filter in the LP560, ProWave 770 and LimeLight eliminates short wavelengths, allowing longer wavelengths to be transmitted to the treatment area. In addition, the wavelength spectrum of the ProWave 770 and the LimeLight can be shifted based on the setting of the control console. Our power control includes a monitoring system to ensure that the desired energy level is delivered. The hand pieces protect the epidermis by regulating the temperature of the hand piece window through the embedded temperature monitor. These hand pieces are available on the Xeo and Solera platforms.

Titan Hand Piece- The Titan hand pieces are designed to produce a sustained pulse of light over a wavelength spectrum tailored to provide heating in the dermis to treat skin laxity (although it is cleared in the United States by the U.S. Food and Drug Administration, or FDA, only for deep dermal heating). The hand piece consists of a custom light source, proprietary wavelength filter, closed-loop power control, sapphire cooling window and embedded temperature monitor, and weighs approximately three pounds. The temperature of the epidermis is controlled by using a sapphire window to provide cooling before, during and after the delivery of energy to the treatment site. We offer two different Titan hand pieces—Titan V and Titan XL.

- Titan V- Titan V has a treatment tip that extends beyond the hand piece housing to provide enhanced visibility of the skin's surface to effectively treat delicate areas such as the skin around the eyes and nose.
- Titan XL- Titan XL, like the Titan V, has a treatment tip that extends beyond the housing for improved visibility. It also has a larger treatment spot size to treat larger body areas faster, such as the arms, abdomen and legs.

The Titan hand pieces can be used on the Xeo and Solera platforms. The Titan hand piece requires a periodic "refilling" process, which includes the replacement of the optical source, after a set number of pulses have been used. This provides us with a source of recurring revenue.

Pearl Hand Piece- The Pearl hand piece, introduced in 2007, is designed to treat fine lines, uneven texture and dyschromia through the application of proprietary YSGG laser technology. This hand piece can safely remove a small portion of the epidermis, while coagulating the remaining epidermis, leading to new collagen growth. The Pearl hand piece consists of a custom monolithic laser source, scanner and power monitoring electronics. The scanner includes multiple scan patterns to allow simple and fast treatments of the face. The hand piece includes an attachment for a smoke evacuator, allowing the practitioner to use one hand during treatment.

Pearl Fractional Hand Piece- The Pearl Fractional hand piece, introduced in 2008, also uses proprietary YSGG technology and is designed to treat wrinkles and deep dermal imperfections (although it is cleared in the United States by the FDA only for skin resurfacing and coagulation). This hand piece penetrates the deep dermis producing a series of microcolumns across the skin, which can result in the removal of damaged tissue and the production of new collagen. The Pearl Fractional hand piece consists of a custom monolithic laser source, scanner and power monitoring electronics. The scanner includes multiple scan patterns to allow simple and fast treatments of the face. The hand piece includes an attachment for a smoke evacuator, allowing the practitioner to use one hand during treatment.

Upgrades

Our Solera and Xeo platforms are multi-application products that are designed to allow our customers to cost-effectively upgrade to our newest technologies, which provides our customers the option to add applications to their system and provides us with a source of recurring revenue. When we introduce a new product, we notify our customers of the upgrade opportunity through a sales call or mailing. In most cases, a field service representative can install the upgrade at the customer site in a matter of hours, which results in very little downtime for practitioners. In some cases, where substantial upgrades are necessary, customers will receive fully-refurbished systems before sending their prior systems back to our headquarters. When customers wish to upgrade from the CoolGlide platform to either a Xeo or a Solera, we provide them with a trade-in value for their CoolGlide and upgrade them to the multi-application platform with the desired applications.

Service

We offer post-warranty services to our customers either through extended service contracts to cover preventive maintenance or replacement parts and labor, or through direct billing for parts and labor. These post-warranty services serve as additional sources of recurring revenue from our installed base.

Titan Hand Piece Refills

Each Titan hand piece is a refillable product, which provides us with a source of recurring revenue from our existing customers.

Fillers and Cosmeceuticals

We distribute BioForm's (a subsidiary of Merz) Radiesse® dermal filler product and Obagi Medical Product, Inc.'s (or Obagi) prescription-based, topical skin health systems (or Cosmeceuticals) to physicians in the Japanese market.

Our Applications and Procedures

Our products are designed to allow the practitioner to select an appropriate combination of energy level, spot size and pulse duration for each treatment. The ability to manipulate the combinations of these parameters allows our customers to treat the broadest range of conditions available with a single energy-based system.

Hair Removal- Our laser technology allows our customers to treat all skin types and hair thicknesses. Our 1064 nm Nd:YAG laser permits energy to safely penetrate through the epidermis of any skin type and into the dermis where the hair follicle is located. Using the universal graphic user interface on our control console, the practitioner sets parameters to deliver therapeutic energy with a large spot size and variable pulse durations, allowing the practitioner to treat fine or coarse hair. Our 1064nm Nd:YAG hand piece allows our customers to treat all skin types, while our ProWave 770 hand piece, with its pulsed light technology, treats the majority of skin types quickly and effectively.

To remove hair using a 1064nm Nd:YAG hand piece, the treatment site on the skin is first cleaned and shaved. The practitioner then applies a thin layer of gel to glide across the skin, and next applies the hand piece directly to the skin to cool the area to be treated and then delivers a laser pulse to the pre-cooled area. To remove hair using the ProWave 770 hand piece, mineral oil is used instead of gel, and cooling is provided by a sapphire window placed directly on the skin, allowing the pulse of light to be applied while the treatment area is being cooled. In the case of both hand pieces, delivery of the energy destroys the hair follicles and prevents hair re-growth. This procedure is then repeated at the next treatment site on the body, and can be done in a gliding motion to increase treatment speed. Patients receive on average three to six treatments. Each treatment can take between five minutes and one hour depending on the size of the area and the condition being treated. On average, there are six to eight weeks between treatments.

Vascular Lesions- Our laser technology allows our customers to treat the widest range of aesthetic vein conditions, including spider and reticular veins and small facial veins. Our CoolGlide and Xeo 1064nm Nd:YAG hand piece's adjustable spot size of 3, 5, 7 or 10 millimeters, or the Excel V 1064 nm and 532 nm hand piece with adjustable spot sizes from 1.5 to 12 mm, allows the practitioner to control treatment depth to target different sized veins. Selection of the appropriate energy level and pulse duration ensures effective treatment of the intended target. Our AcuTip 500 hand piece, with its 6 millimeter spot size, uses pulsed-light technology and is designed for the treatment of facial vessels.

The vein treatment procedure when using the 1064nm Nd:YAG hand piece is performed in a substantially similar manner to the laser hair removal procedure. The laser hand piece is used to cool the treatment area both before and after the laser pulse has been applied. With the Excel V hand piece the cooling can be performed pre, during and post delivery of the laser pulse. With the AcuTip 500 hand piece, the pulse of light is delivered while the treatment area is being cooled with the sapphire tip. The delivered energy damages the vein and, over time, it is absorbed by the body. Patients receive on average between one and six treatments, with six weeks or longer between treatments.

Skin Rejuvenation- Our laser and light-based technologies allow our customers to perform non-invasive and minimally-invasive treatments that reduce redness, pore size, fine lines and laxity, improve skin texture, and treat other aesthetic conditions. Our products are each designed to minimize the risk of damage to the surrounding tissue.

Texture; Lines and Wrinkles- When using a 1064nm Nd: YAG laser to improve skin texture, reduce pore size and treat fine lines, cooling is not applied and the hand piece is held directly above the skin. A large number of pulses are directed at the treatment site, repeatedly covering an area, such as the cheek. By delivering many pulses of laser light to a treatment area, a gentle heating of the dermis occurs and collagen growth is stimulated to rejuvenate the skin and reduce wrinkles. Patients typically receive four to six treatments for this procedure. The treatment typically takes less than a half hour and there are typically two to four weeks between treatments.

When treating texture and fine lines with a Pearl hand piece, the hand piece is held at a controlled distance from the skin and the scanner delivers a preset pattern of spots to the treatment area. Cooling is not applied to the epidermis during the treatment. The energy delivered by the hand piece ablates a portion of the epidermis while leaving a coagulated portion that will gently peel off over the course of a few days. Heat is also delivered into the dermis which can result in the production of new collagen. Treatment of the full face can usually be performed in 15 to 30 minutes. Patients receive on average between one and three treatments at monthly intervals.

When treating wrinkles and deep dermal imperfections with a Pearl Fractional hand piece, the hand piece is held at a controlled distance from the skin and the scanner delivers a preset pattern of spots to the treatment area. Cooling is not applied to the epidermis during the treatment. The energy delivered by the hand piece penetrates the deep dermis producing a series of microcolumns across the skin, which can result in the removal of damaged tissue and the production of new collagen. Treatment of the full face can usually be performed in less than an hour. Patients receive on average between one and three treatments at monthly intervals.

Our CE Mark allows us to market Pearl Fractional in the European Union, Australia and certain other countries outside the United States for the treatment of wrinkles and deep dermal imperfections. However, in the United States we have a 510(k) clearance for only skin resurfacing and coagulation.

Toenail Fungus- In addition to performing skin rejuvenation, we have a CE Mark approval for the GenesisPlus that allows us to market it in the European Union and certain other countries outside the United States for the treatment of onychomycosis (or toenail fungus). We have submitted a 501(k) application to the FDA to market this product in the United States for the treatment of toenail fungus and do not currently have approval. Tiny pulses of light from an Nd: YAG laser pass through the toenail to the fungus underneath, which is irradiated without any damage to the surrounding nail or skin. The GenesisPlus has two aiming beams that facilitate consistent treatments by maintaining the correct distance of the hand piece to the skin. In addition, during the treatment an integrated sensor is used to actively monitor the temperature of the treatment area.

Dyschromia- Our pulsed-light technologies allow our customers to safely and effectively treat red and brown dyschromia, which is skin discoloration, pigmented lesions and rosacea. The practitioner delivers a narrow spectrum of light to the surface of the skin through our LP560 or LimeLight hand pieces. These hand pieces include one of our proprietary wavelength filters, which reduce the energy level required for therapeutic effect and minimize the risk of skin injury.

In treating pigmented lesions with a pulsed-light technology, the hand piece is placed directly on the skin and then the light pulse is triggered. The cells forming the pigmented lesion absorb the light energy, darken and then flake off over the course of two to three weeks. Several treatments may be required to completely remove the lesion. The treatment takes a few minutes per area treated and there are typically three to four weeks between treatments.

The 532 nm wavelength green laser option on the Excel V can also be used to treat pigmented lesions in substantially the same way as described above with the pulsed light devices.

Practitioners can also treat dyschromia and other skin conditions with our Pearl hand piece. During these treatments, the heat delivered by the Pearl hand piece will remove the outer layer of the epidermis while coagulating a portion of the epidermis. That coagulated portion will gently peel off over the course of a few days, revealing a new layer of skin underneath. Treatment of the full face can usually be performed in 15 to 30 minutes. Patients receive on average between one and three treatments at monthly intervals.

Skin Laxity-Our Titan technology allows our customers to use deep dermal heating to tighten lax skin. The practitioner delivers a spectrum of light to the skin through our Titan hand piece. This hand piece includes our proprietary light source and wavelength filter which tailors the delivered spectrum of light to provide heating at the desired depth in the skin.

In treating skin laxity, the hand piece is placed directly on the skin and then the light pulse is triggered. A sustained pulse causes significant heating in the dermis. This heating can cause immediate collagen contraction while also stimulating long-term collagen re-growth. Several treatments may be required to obtain the desired degree of tightening of the skin. The treatment of a full face can take over an hour and there are typically four weeks between treatments.

Our CE Mark allows us to market the Titan in the European Union, Australia and certain other countries outside the United States for the treatment of wrinkles through skin tightening. However, in the United States we have a 510(k) clearance for only deep dermal heating.

Sales and Marketing

In the United States we market and sell our products primarily through a direct sales organization. Generally, each direct sales employee is assigned a specific territory. As of December 31, 2010, we had a U.S. direct sales force of 19 employees. We internally manage our U.S. and Canadian sales organization as one North American sales region with 27 territories as of December 31, 2010. In addition to direct sales employees, we have a distribution relationship with PSS World Medical that operates medical supply distribution service centers with over 700 sales representatives serving physician offices throughout the United States. Revenue from PSS was \$2.6 million in 2010, \$3.8 million in 2009, and \$12.1 million in 2008.

International sales are generally made through a direct international sales force of 26 employees, as well as a worldwide distributor network in over 35 countries as of December 31, 2010. As of December 31, 2010, we had direct sales offices in Australia, Canada, France, Japan, Spain, Switzerland and the United Kingdom. Our international revenue as a percentage of total revenue represented 64% in 2010, 61% in 2009, and 50% in 2008.

We also sell certain items like Titan hand piece refills and marketing brochures via the internet.

Although specific customer requirements can vary depending on applications, customers generally demand quality, performance, ease of use, and high productivity in relation to the cost of ownership. We have responded to these customer demands by introducing new products focused on these requirements in the markets we serve. Specifically, we believe that we introduce new products and applications that are innovative, address the specific aesthetic procedures in demand, and are upgradeable on our customers' existing systems. In addition, we provide attractive upgrade pricing to new product families and are responsive to our customers' financing preferences. To increase market penetration, in addition to marketing to the core specialties of plastic surgeons and dermatologists, we also market to the non-core aesthetic practices consisting of gynecologists, primary care physicians, family practitioners, physicians offering aesthetic treatments in non-medical offices and other qualified practitioners.

We seek to establish strong ongoing relationships with our customers through the upgradeability of our products, sales of extended service contracts, the refilling of Titan hand pieces, ongoing training and support, and distributing (in Japan only) a dermal filler product. We primarily target our marketing efforts to practitioners through office visits, workshops, trade shows, webinars and trade journals. We also market to potential patients through brochures, workshops and our website. In addition, we offer clinical forums with recognized expert panelists to promote advanced treatment techniques using our products to further enhance customer loyalty and uncover new sales opportunities.

Competition

Our industry is subject to intense competition. Our products compete against conventional non-energy-based treatments, such as electrolysis, Botox and collagen injections, chemical peels, microdermabrasion and sclerotherapy. Our products also compete against laser and light-based products offered by public companies, such as Cynosure, Elen (in Italy), Iridex, Palomar, Solta and Syneron, as well as private companies, including, Alma, Lumenis, Sciton and several other companies.

Competition among providers of laser and other energy-based devices for the aesthetic market is characterized by extensive research efforts and innovative technology. While we attempt to protect our products through patents and other intellectual property rights, there are few barriers to entry that would prevent new entrants or existing competitors from developing products that would compete directly with ours. There are many companies, both public and private, that are developing innovative devices that use both energy-based and alternative technologies. Some of these competitors have greater resources than we do or product applications for certain sub-markets in which we do not participate. Additional competitors may enter the market, and we are likely to compete with new companies in the future. To compete effectively, we have to demonstrate that our products are attractive alternatives to other devices and treatments by differentiating our products on the basis of performance, brand name, service and price. We have encountered, and expect to continue to encounter, potential customers who, due to existing relationships with our competitors, are committed to, or prefer the products offered by these competitors. Competitive pressures may result in price reductions and reduced margins for our products.

Research and Development

Our research and development group develops new products and applications and builds clinical support to address unmet or underserved market needs. As of December 31, 2010, our research and development activities were conducted by a staff of 24 employees with a broad base of experience in lasers, optoelectronics, software and other fields. We have developed relationships with outside contract engineering and design consultants, giving our team additional technical and creative breadth. We work closely with thought leaders and customers, to understand unmet needs and emerging applications in aesthetic medicine. Research and development expenses were approximately \$7.0 million in 2010, \$6.8 million in 2009 and \$7.6 million in 2008.

Service and Support

Our products are engineered to enable quick and efficient service and support. There are several separate components of our products, each of which can easily be removed and replaced. We believe that quick and effective delivery of service is important to our customers. As of December 31, 2010, we had a 31-person global service department. Internationally, we provide direct service support through our Australia, Canada, France, Japan, Spain and Switzerland offices, and also through the network of distributors in over 35 countries and third-party service providers. We historically have provided a standard one-year or two-year warranty coverage on our systems. We have a standard oneyear warranty on all systems. We provide initial warranties on our products to cover parts and service and offer extended service plans that vary by the type of product and the level of service desired. Our standard warranty on system consoles covers parts and service for a standard period of one year. From time to time, we also have promotions whereby we include a post-warranty service contract with the sale of our products. Customers are notified before their initial warranty expires and are able to choose from two different extended service plans covering preventative maintenance or replacement parts and labor. In the event a customer does not purchase an extended service plan, we will offer to service the customer's system and charge the customer for time and materials. Our Titan hand pieces generally include a warranty for a set number of shots instead of for a period of time. We have invested substantial financial and management resources to develop a worldwide infrastructure to meet the service needs of our customers worldwide.

Manufacturing

We manufacture our products with components and subassemblies supplied by vendors. We assemble and test each of our products at our Brisbane, California facility. Quality control, cost reduction and inventory management are top priorities of our manufacturing operations.

We purchase certain components and subassemblies from a limited number of suppliers. We have flexibility with our suppliers to adjust the number of components and subassemblies as well as the delivery schedules. The forecasts we use are based on historical demands and sales projections. Lead times for components and subassemblies may vary significantly depending on the size of the order, time required to fabricate and test the components or subassemblies, specific supplier requirements and current market demand for the components and subassemblies. We reduce the potential for disruption of supply by maintaining sufficient inventories and identifying additional suppliers. The time required to qualify new suppliers for some components, or to redesign them, could cause delays in our manufacturing. To date, we have not experienced significant delays in obtaining any of our components or subassemblies.

We use small quantities of common cleaning products in our manufacturing operations, which are lawfully disposed of through a normal waste management program. We do not forecast any material costs due to compliance with environmental laws or regulations.

We are required to manufacture our products in compliance with the FDA's Quality System Regulation, or QSR. The QSR covers the methods and documentation of the design, testing, control, manufacturing, labeling, quality assurance, packaging, storage and shipping of our products. The FDA enforces the QSR through periodic unannounced inspections. We had a full quality system audit in 2008 and an FDA audit of compliance with laser performance standards in 2010 for our single manufacturing facility located in Brisbane, CA. There were no significant findings as a result of this audit and our responses have been accepted by the FDA. Our failure to maintain compliance with the QSR requirements could result in the shut down of our manufacturing operations and the recall of our products, which would have a material adverse effect on business. In the event that one of our suppliers fails to maintain compliance with our quality requirements, we may have to qualify a new supplier and could experience manufacturing delays as a result. We have opted to maintain quality assurance and quality management certifications to enable us to market our products in the United States, the member states of the European Union, the European Free Trade Association and countries which have entered into Mutual Recognition Agreements with the European Union. Our manufacturing facility is ISO 13485 certified.

Patents and Proprietary Technology

We rely on a combination of patent, copyright, trademark and trade secret laws, and non-disclosure, confidentiality and invention assignment agreements to protect our intellectual property rights. As of December 31, 2010, we had 17 issued U.S. patents and 24 pending U.S. patent applications. Acutip 500, Cutera, CoolGlide, CoolGlide Excel, Limelight, Pearl, Pearl Fractional, ProWave 770, Solera, Solera Titan, Titan and Xeo are only some of the trademarks and/or service marks of Cutera in the U.S. and other countries. We have trademark rights to these names and others in the United States and certain other countries. We intend to file for additional patents and trademarks to continue to strengthen our intellectual property rights.

We license certain patents from Palomar and pay ongoing royalties based on sales of applicable hair-removal products. The royalty rate on these products ranges from 3.75% to 7.50% of revenue. The patents are set to expire in February 2013 and February 2015. Our revenue from systems that do not include hair-removal capabilities (such as our Solera Titan, Xeo SA, GenesisPlus and Excel V); and other revenue from service contracts, Titan refills, Fillers and cosmeceuticals, are not subject to these royalties. In addition, in 2006 we capitalized \$1.2 million as an intangible asset representing the ongoing license for these patents, which is being amortized on a straight-line basis over their expected useful life of 9-10 years. We also have a technology sublicense purchased in 2002, which is being amortized on a straight-line basis over its expected useful life of 10 years.

Our employees and technical consultants are required to execute confidentiality agreements in connection with their employment and consulting relationships with us. We also require them to agree to disclose and assign to us all inventions conceived in connection with the relationship. We cannot provide any assurance that employees and consultants will abide by the confidentiality or assignability terms of their agreements. Despite measures taken to protect our intellectual property, unauthorized parties may copy aspects of our products or obtain and use information that we regard as proprietary.

Government Regulation

Our products are medical devices subject to extensive and rigorous regulation by the U.S. Food and Drug Administration, as well as other regulatory bodies. FDA regulations govern the following activities that we perform and will continue to perform to ensure that medical products distributed domestically or exported internationally are safe and effective for their intended uses:

- Product design and development;
- Product testing;
- Product manufacturing:
- Product safety;
- Product labeling;
- Product storage;
- Recordkeeping;
- Pre-market clearance or approval;
- Advertising and promotion;
- Production; and
- Product sales and distribution.

FDA's Pre-market Clearance and Approval Requirements

Unless an exemption applies, each medical device we wish to commercially distribute in the United States will require either prior 510(k) clearance or pre-market approval from the FDA. The FDA classifies medical devices into one of three classes. Devices deemed to pose lower risks are placed in either class I or II, which requires the manufacturer to submit to the FDA a pre-market notification requesting permission to commercially distribute the device. This process is generally known as 510(k) clearance. Some low risk devices are exempted from this requirement. Devices deemed by the FDA to pose the greatest risk, such as life-sustaining, life-supporting or implantable devices, or devices deemed not substantially equivalent to a previously cleared 510(k) device, are placed in class III, requiring pre-market approval. All of our current products are class II devices.

510(k) Clearance Pathway

When a 510(k) clearance is required, we must submit a pre-market notification demonstrating that our proposed device is substantially equivalent to a previously cleared 510(k) device or a device that was in commercial distribution before May 28, 1976 for which the FDA has not yet called for the submission of Pre-Market Approval, or PMA, applications. By regulation, the FDA is required to clear or deny a 510(k), pre-market notification within 90 days of submission of the application. As a practical matter, clearance often takes significantly longer. The FDA may require further information, including clinical data, to make a determination regarding substantial equivalence. Laser devices used for aesthetic procedures, such as hair removal, have generally qualified for clearance under 510(k) procedures.

The following table details the indications for which we received a 510(k) clearance for our products and when these clearances were received.

FDA Marketing Clearances:	Date Received:
Laser-based products:	
- treatment of vascular lesions	June 1999
- hair removal	March 2000
- permanent hair reduction	January 2001
- treatment of benign pigmented lesions and pseudofolliculitis barbae, commonly referred to	
as razor bumps, and for the reduction of red pigmentation in scars	June 2002
- treatment of wrinkles	October 2002
Pulsed-light technologies:	
- treatment of pigmented lesions	March 2003
- hair removal and vascular treatments	March 2005
Infrared Titan technology for deep dermal heating for the temporary relief of minor muscle and joint pain and for the temporary increase in local circulation where applied	February 2004
Solera tabletop console:	
- for use with the Titan hand piece	October 2004
- for use with our pulsed-light hand pieces	January 2005
Pearl product for the treatment of wrinkles	March 2007
Pearl Fractional product for skin resurfacing and coagulation	August 2008

Pre-Market Approval (PMA) Pathway

A PMA must be submitted to the FDA if the device cannot be cleared through the 510(k) process. A PMA must be supported by extensive data, including but not limited to, technical, preclinical, clinical trials, manufacturing and labeling to demonstrate to the FDA's satisfaction the safety and effectiveness of the device. No device that we have developed to date has required pre-market approval, although development of future devices or indications may require pre-market approval.

Product Modifications

We have modified aspects of our products since receiving regulatory clearance, but we believe that new 510(k) clearances are not required for these modifications. After a device receives 510(k) clearance or a PMA, any modification that could significantly affect its safety or effectiveness, or that would constitute a major change in its intended use, will require a new clearance or approval. The FDA requires each manufacturer to make this determination initially, but the FDA can review any such decision and can disagree with a manufacturer's determination. If the FDA disagrees with our determination not to seek a new 510(k) clearance or PMA, the FDA may retroactively require us to seek 510(k) clearance or pre-market approval. The FDA could also require us to cease marketing and distribution and/or recall the modified device until 510(k) clearance or pre-market approval is obtained. Also, in these circumstances, we may be subject to significant regulatory fines or penalties.

Clinical Trials

When FDA approval of a class I, class II or class III device requires human clinical trials, and if the device presents a "significant risk," as defined by the FDA, to human health, the device sponsor is required to file an Investigational Device Exemption, or IDE, application with the FDA and obtain IDE approval prior to commencing the human clinical trial. If the device is considered a "non-significant" risk, IDE submission to the FDA is not required. Instead, only approval from the Institutional Review Board, or IRB, overseeing the clinical trial is required. Human clinical studies are generally required in connection with approval of class III devices and may be required for class I and II devices. The IDE application must be supported by appropriate data, such as animal and laboratory testing results, showing that it is safe to test the device in humans and that the testing protocol is scientifically sound. The IDE must be approved in advance by the FDA for a specified number of patients. Clinical trials for a significant risk device may begin once the application is reviewed and cleared by the FDA and the appropriate institutional review boards at the clinical trial sites. Future clinical trials of our products may require that we submit and obtain clearance of an IDE from the FDA prior to commencing clinical trials. The FDA, and the IRB at each institution at which a clinical trial is being performed, may suspend a clinical trial at any time for various reasons, including a belief that the subjects are being exposed to an unacceptable health risk.

Pervasive and Continuing Regulation

After a device is placed on the market, numerous regulatory requirements apply. These include:

- Quality system regulations, which require manufacturers, including third-party manufacturers, to follow stringent design, testing, control, documentation and other quality assurance procedures during all aspects of the manufacturing process;
- Labeling regulations and FDA prohibitions against the promotion of products for un-cleared, unapproved or "off-label" uses;
- Medical device reporting regulations, which require that manufacturers report to the FDA if their device
 may have caused or contributed to a death or serious injury or malfunctioned in a way that would likely
 cause or contribute to a death or serious injury if the malfunction were to recur; and
- Post-market surveillance regulations, which apply when necessary to protect the public health or to provide additional safety and effectiveness data for the device.

The FDA has broad post-market and regulatory enforcement powers. We are subject to unannounced inspections by the FDA and the Food and Drug Branch of the California Department of Health Services, or CDHS, to determine our compliance with the QSR and other regulations, and these inspections may include the manufacturing facilities of our subcontractors. In the past, our prior facility has been inspected, and observations were noted. There were no findings that involved a material violation of regulatory requirements. Our responses to these observations have been accepted by the FDA and CDHS, and we believe that we are in substantial compliance with the QSR. Our current manufacturing facility has been inspected by the FDA and the CDHS. The FDA and the CDHS noted observations, but there were no findings that involved a material violation of regulatory requirements. Our responses to those observations have been accepted by the FDA and CDHS.

We are also regulated under the Radiation Control for Health and Safety Act, which requires laser products to comply with performance standards, including design and operation requirements, and manufacturers to certify in product labeling and in reports to the FDA that their products comply with all such standards. The law also requires laser manufacturers to file new product and annual reports, maintain manufacturing, testing and sales records, and report product defects. Various warning labels must be affixed and certain protective devices installed, depending on the class of the product.

Failure to comply with applicable regulatory requirements can result in enforcement action by the FDA, which may include any of the following sanctions:

- Warning letters, fines, injunctions, consent decrees and civil penalties;
- Repair, replacement, recall or seizure of our products;
- Operating restrictions or partial suspension or total shutdown of production;
- Refusing our requests for 510(k) clearance or pre-market approval of new products, new intended uses, or modifications to existing products;

- Withdrawing 510(k) clearance or pre-market approvals that have already been granted; and
- Criminal prosecution.

The FDA also has the authority to require us to repair, replace or refund the cost of any medical device that we have manufactured or distributed. If any of these events were to occur, they could have a material adverse effect on our business.

We are also subject to a wide range of federal, state and local laws and regulations, including those related to the environment, health and safety, land use and quality assurance. We believe that compliance with these laws and regulations as currently in effect will not have a material adverse effect on our capital expenditures, earnings and competitive and financial position.

International

International sales of medical devices are subject to foreign governmental regulations, which vary substantially from country to country. The time required to obtain clearance or approval by a foreign country may be longer or shorter than that required for FDA clearance or approval, and the requirements may be different.

The primary regulatory environment in Europe is that of the European Union, which consists of a number of countries encompassing most of the major countries in Europe. The member states of the European Free Trade Association have voluntarily adopted laws and regulations that mirror those of the European Union with respect to medical devices. Other countries, such as Switzerland, have entered into Mutual Recognition Agreements and allow the marketing of medical devices that meet European Union requirements. The European Union has adopted numerous directives and European Standardization Committees have promulgated voluntary standards regulating the design, manufacture, clinical trials, labeling and adverse event reporting for medical devices. Devices that comply with the requirements of a relevant directive will be entitled to bear CE conformity marking, indicating that the device conforms with the essential requirements of the applicable directives and, accordingly, can be commercially distributed throughout the member states of the European Union, the member states of the European Free Trade Association and countries which have entered into a Mutual Recognition Agreement. The method of assessing conformity varies depending on the type and class of the product, but normally involves a combination of self-assessment by the manufacturer and a third-party assessment by a Notified Body, an independent and neutral institution appointed by a country to conduct the conformity assessment. This third-party assessment may consist of an audit of the manufacturer's quality system and specific testing of the manufacturer's device. An assessment by a Notified Body in one member state of the European Union, the European Free Trade Association or one country which has entered into a Mutual Recognition Agreement is required in order for a manufacturer to commercially distribute the product throughout these countries. ISO 9001 and ISO 13845 certification are voluntary harmonized standards. Compliance establishes the presumption of conformity with the essential requirements for a CE Marking. In February 2000, our facility was awarded the ISO 9001 and EN 46001 certification. In March 2003, we received our ISO 9001 updated certification (ISO 9001:2000) as well as our certification for ISO 13485:1996 which replaced our EN 46001 certification. In March 2004, we received our ISO 13485:2003 certification, which is the most current ISO certification for medical device companies, and in March 2006, March 2010, and February 2011 we passed our ISO 13485 recertification audits.

Employees

As of December 31, 2010, we had 187 employees, compared to 186 employees as of December 31, 2009. Of the 187 employees at December 31, 2010, 77 were in sales and marketing, 33 in manufacturing operations, 31 in technical service, 24 in research and development and 22 in general and administrative. We believe that our future success will depend in part on our continued ability to attract, hire and retain qualified personnel. None of our employees are represented by a labor union, and we believe our employee relations are good.

Available Information

We are subject to the reporting requirements under the Securities Exchange Act of 1934. Consequently, we are required to file reports and information with the Securities and Exchange Commission, or SEC, including reports on the following forms: annual report on Form 10-K, quarterly reports on Form 10-Q, current reports on Form 8-K, and amendments to those reports filed or furnished pursuant to Section 13(a) or 15(d) of the Securities Exchange Act of 1934. These reports and other information concerning the company may be accessed through the SEC's website at http://www.sec.gov. Such filings, as well as our charters for our Audit and Compensation Committees and our Code of Ethics are available on our website at http://www.cutera.com. In the event that we grant a waiver under our Code of Ethics to any of our officers and directors, we will publish it on our website.

ITEM 1A. RISK FACTORS

The recent earthquake and tsunami, and other collateral events, in Japan may adversely affect the demand for our products and services in the Japanese market, which may cause a decline in revenues and negatively affect our operating results.

We have two direct sales offices in Japan and generate revenue from the sale of systems, upgrades, Titan refills, fillers, cosmeceuticals and services. Revenue sourced from the Japanese market was approximately \$13.6 million in 2010, \$9.6 million in 2009 and \$10.9 million in 2008. In 2010, our Japanese sourced revenue represented 26% of our world wide revenue and we experienced growth from all of our product categories, including Fillers and Cosmeceuticals due to the commencement of the distribution of cosmeceutical products in 2010.

The recent earthquake and tsunami in Japan, and other collateral events, including, among others, the catastrophic loss of lives, businesses, infrastructure, and delays in transportation, may have a direct negative impact on us or an indirect impact on us by affecting our employees, customers, or the overall economy in Japan and may reduce the demand for our products and services. As a result, these events could cause a decline in our revenue in Japan and our results of operations could be materially and adversely affected.

In 2010, our U.S. revenue decreased by approximately 8%, compared to the same period in 2009. Our U.S. revenue is significantly below the pre-2009 level due in part to customers purchasing fewer applications and declining average selling prices (or ASPs). In addition, we experienced a temporary decline in our Titan refill revenue in 2010, compared to 2009, due to a voluntary recall of certain Titan XL hand pieces. All customers that had a Titan XL hand pieces subject to the recall were provided with fully refilled hand pieces, which delayed their purchase of a refill. If our U.S. revenue does not improve, it could have a material adverse effect on our total revenue, profitability, employee retention and stock price.

In 2010, our U.S. revenue decreased by approximately 8%, compared to 2009. Our U.S. revenue is significantly below the pre-2009 level due to several factors, some of which are:

- Our Product and Upgrade ASPs in 2010 were lower than the pre-2010 levels as a result of customers purchasing fewer applications for systems and lower pricing resulting from competitive discounting pressures.
- Historically, we have introduced a new product every year since 2000, which typically resulted in increased revenue. However, in 2009 and until August 2010, we did not have a new product. In 2010 we launched GenesisPlus and in February 2011, we introduced our Excel V laser system a unique vascular work station designed specifically for the core-market of Dermatologists and Plastic Surgeons. However, even though we have introduced these new products, there can be no assurance that they will translate into increased revenue in the U.S. in 2011.
- We experienced a temporary decline in our Titan refill revenue in 2010, compared to 2009, due to a voluntary recall of certain Titan XL hand pieces. All customers that had a Titan XL hand piece subject to the recall were provided with fully refilled hand pieces, which delayed their purchase of a refill.

If our U.S. revenue does not improve, it could have a material adverse effect on our total revenue, profitability, employee retention and stock price.

We rely heavily on our sales professionals to market and sell our products worldwide. If we are unable to hire, effectively train, manage, improve the productivity of, and retain the sales professionals, our business will be harmed, which would impair our future revenue and profitability.

Our success largely depends on our ability to hire, train, manage and improve the productivity levels of our sales professionals worldwide. Because of our focus on the non-core market in the past, several of our sales professionals do not have established relationships with core market physicians (Dermatologists and Plastic Surgeons) or where those relationships exist, they are not very strong. In addition, we have lost some of our sales professionals in response to the decline in their earnings resulting from the decreases in their commissions.

We have selectively hired, and continue to do so, new sales professionals in key territories to fill vacant positions. For example, in December 2010, Michael Poole joined us as Vice President of North American Sales, which allowed our previous Vice President of North American Sales to return to Japan in an expanded role to lead our Pacific Rim operations. We have been training our existing, and recently recruited, sales professionals to better understand our product technology and how it can be positioned against our competitors' products. These initiatives are intended to improve our revenue and profitability.

Although Mr. Poole has over 17 years of a broad range of sales experience and was employed by us from 2004 to 2008, Mr. Poole has limited prior experience in managing a large sales force. Measures we implement in an effort to retain, train and manage our sales professionals, strengthen their relationships with core market physicians, and improve their productivity may not be successful and may instead contribute to instability in our operations, additional departures from our sales organization, or further reduce our revenue and harm to our business.

If our revenue does not improve from the 2010 level, or if our cost of revenue and/or operating expenses increase, our gross margins and operating margins may be adversely impacted, our loss from operations will increase, and our cash used in operating activities will increase, which could reduce our assets and have a material adverse effect on our stock price.

Our gross margin in 2010 was 57%, compared with 59% in 2009 and 61% in 2008. Our gross margin is impacted by the revenue that we generate and the costs incurred to generate the revenue. Our future revenue may be adversely affected by a number of factors including, the competitive market environment in which we operate, which may result in a decrease in the number of units sold, a decrease in the number of applications per system purchased by customers, a decrease in the average selling prices achieved for our product sales, or a shift in our product mix. Our cost of revenue may also be adversely impacted by various factors such as obsolescence of our inventory, increased expenses associated with repairing defective products covered by our warranty program, and utilization of our relatively fixed manufacturing costs.

In response to reduced revenue, in 2009 we reduced our expense levels in various departments including sales, marketing, customer relations, manufacturing, and service departments. We implemented three reduction-in-force (or RIF) programs in 2009, specifically targeting positions that were directly impacted by the recession. Even though we implemented the restructurings, we had a significant loss from operations and used significant cash in our operations in 2010.

If our revenue does not improve from the 2010 level, or if our cost of revenue increases, or if we are not able to reduce expenses in the event of a decline in revenue, we may continue to generate losses from operations and use cash, which could reduce our assets and have a material adverse effect on our operations and stock price.

Demand for our products in any of our markets could be weakened by several factors, including:

- Our ability to develop and market our products to the core market specialties of dermatologists and plastic surgeons;
- Poor financial performance of market segments that try introducing aesthetic procedures to their businesses;
- The inability to differentiate our products from those of our competitors;
- Reduced patient demand for elective aesthetic procedures:
- Failure to build and maintain relationships with opinion leaders within the various market segments;

- An increase in malpractice lawsuits that result in higher insurance costs; and
- The lack of credit financing for some of our potential customers.

If we do not achieve anticipated demand for our products, it could have a material adverse effect on our total revenue, profitability, employee retention and stock price.

The aesthetic equipment market is characterized by rapid innovation. To compete effectively, we must develop and/or acquire new products, market them successfully, and identify new markets for our technology.

We have created products to apply our technology to hair removal, treatment of veins and skin rejuvenation, including the treating of diffuse redness, skin laxity, fine lines, wrinkles, skin texture, pore size and pigmented lesions. Currently, these applications represent the majority of offered laser and light-based aesthetic procedures. We have recently started distributing topical skin creams and dermal fillers in the Japanese market. To grow in the future, we must develop and acquire new and innovative aesthetic products and applications, identify new markets, and successfully launch the newly acquired or developed product offerings.

To successfully expand our product offerings, we must, among other things:

- Develop and acquire new products that either add to or significantly improve our current product offerings;
- Convince our existing and prospective customers that our product offerings would be an attractive revenue-generating addition to their practice;
- Sell our product offerings to a broad customer base;
- Identify new markets and alternative applications for our technology;
- Protect our existing and future products with defensible intellectual property; and
- Satisfy and maintain all regulatory requirements for commercialization.

Except for 2009, we have introduced a new product every year since 2000. In February 2011, we announced the release of our Excel V laser system, a unique vascular work station designed specifically for the core-market of Dermatologists and Plastic Surgeons. In 2010, we launched GenesisPlus, a laser specifically created for the aesthetic treatment of toes and feet. Historically, product introductions have been a significant component of our financial performance. To be successful in the aesthetics industry, we need to continue to innovate. Our business strategy has therefore been based, in part; on our expectation that we will continue to increase our product offerings. We need to continue to devote substantial research and development resources to make new product introductions, which can be costly and time consuming to our organization.

In January 2011, we announced the appointment of Len DeBenedictis as Chief Technology Officer to lead our research and product development efforts. Our current Vice President of Research and Development will report to Mr. DeBenedictis. Although Mr. DeBenedictis has over 20 years of laser and light-based industry experience and an outstanding background to lead our research and product development efforts, there is no guarantee that we will be able to continue our trend of regular new product introductions or that such management change will result in an improved research and development organization. Also, we may need additional research and development resources to make new product introductions, which may be more costly and time consuming to our organization.

We also believe that, to increase revenue from sales of new products and related upgrades, we need to continue to develop our clinical support, further expand and nurture relationships with industry thought leaders and increase market awareness of the benefits of our new products. However, even with a significant investment in research and development, we may be unable to continue to develop, acquire or effectively launch and market new products and technologies regularly, or at all. If we fail to successfully commercialize new products, our business may be harmed.

While we attempt to protect our products through patents and other intellectual property, there are few barriers to entry that would prevent new entrants or existing competitors from developing products that compete directly with ours. For example, while our CoolGlide product was the first long-pulse Nd:YAG, or long wavelength, laser system cleared by the FDA for permanent hair reduction on all skin types, competitors have subsequently introduced systems that utilize Nd:YAG lasers, and received FDA clearances to market these products as treating all skin types. We expect that any competitive advantage we may enjoy from other current and future innovations, such as combining multiple hand pieces in a single system to perform a variety of applications, may diminish over time as companies successfully respond to our, or create their own, innovations. Consequently, we believe that we will have to continuously innovate and improve our products and technology to compete successfully. If we are unable to innovate successfully, our products could become obsolete and our revenue could decline as our customers and prospects purchase our competitors' products.

Our ability to effectively compete and generate additional revenue from new and existing products depend upon our ability to distinguish our company and our products from our competitors and their products, and to develop and effectively market new and existing products. Our success is dependent on many factors, including the following:

- Speed of new and innovative product development;
- Effective strategy and execution of new product launches;
- Identify and develop clinical support for new indications of our existing products;
- Product performance;
- Product pricing;
- Quality of customer support:
- Development of successful distribution channels, both domestically and internationally; and
- Intellectual property protection.

To compete effectively, we have to demonstrate that our new and existing products are attractive alternatives to other devices and treatments, by differentiating our products on the basis of such factors as innovation, performance, brand name, service, and price. This is difficult to do, especially in a crowded aesthetic market. Some of our competitors have newer or different products and more established customer relationships than we do, which could inhibit our market penetration efforts. For example, we have encountered, and expect to continue to encounter, situations where, due to pre-existing relationships, potential customers decided to purchase additional products from our competitors. Potential customers also may need to recoup the cost of products that they have already purchased from our competitors and may decide not to purchase our products, or to delay such purchases.

If we are unable to increase our market penetration or compete effectively, our revenue and profitability will be adversely impacted.

We compete against companies that offer alternative solutions to our products, or have greater resources, a larger installed base of customers and broader product offerings than ours. If we are not able to effectively compete with these companies, it may harm our business.

Our industry is subject to intense competition. Our products compete against similar products offered by public companies, such as Cynosure, Elen (in Italy), Iridex, Palomar, Solta, and Syneron and as well as private companies such as Alma, Lumenis, Sciton and several other companies. Recently, there has been consolidation in the aesthetic industry leading to companies combining their resources. For example, Solta (previously Thermage) acquired Reliant in December 2008 and Aesthera in February 2010; and Syneron acquired Candela in September 2009. We are likely to compete with new companies in the future. Competition with these companies could result in reduced selling prices, reduced profit margins and loss of market share, any of which would harm our business, financial condition and results of operations.

The energy-based aesthetic market faces competition from non energy-based medical products, such as Botox, an injectable compound used to reduce wrinkles, and collagen injections. Other alternatives to the use of our products include electrolysis, a procedure involving the application of electric current to eliminate hair follicles, and chemical peels. We may also face competition from manufacturers of pharmaceutical and other products that have not yet been developed.

If there is not sufficient consumer demand for the procedures performed with our products, practitioner demand for our products could be inhibited, resulting in unfavorable operating results and reduced growth potential.

Continued expansion of the global market for laser and light-based aesthetic procedures is a material assumption of our business strategy. Most procedures performed using our products are elective procedures not reimbursable through government or private health insurance, with the costs borne by the patient. The decision to utilize our products may therefore be influenced by a number of factors, including:

- Consumer disposable income and access to consumer credit, which as a result of the unstable economy, may have been significantly impacted;
- The cost of procedures performed using our products;
- The cost, safety and effectiveness of alternative treatments, including treatments which are not based upon laser or light-based technologies and treatments which use pharmaceutical products;
- The success of our sales and marketing efforts; and
- The education of our customers and patients on the benefits and uses of our products, compared to competitors' products and technologies.

If, as a result of these factors, there is not sufficient demand for the procedures performed with our products, practitioner demand for our products could be reduced, which could have a material adverse effect on our business, financial condition, revenue and result of operations.

Any defects in the design, material or workmanship of our products may not be discovered prior to shipment to customers, which could materially increase our expenses, adversely impact profitability and harm our business.

The design of our products is complex. To manufacture them successfully, we must procure quality components and employ individuals with a significant degree of technical expertise. If our designs are defective, or the material components used in our products are subject to wearing out, or if suppliers fail to deliver components to specification, or if our employees fail to properly assemble, test and package our products, the reliability and performance of our products will be adversely impacted. As an example, in 2010, we incurred significant expenses for the voluntary recall of our Titan XL hand pieces.

If our products contain defects that cannot be repaired easily, inexpensively, or on a timely basis, we may experience:

- Damage to our brand reputation;
- Loss of customer orders and delay in order fulfillment;
- Increased costs due to product repair or replacement;
- Inability to attract new customers;
- Diversion of resources from our manufacturing and research and development departments into our service department; and
- Legal action.

The occurrence of any one or more of the foregoing could materially increase expenses, adversely impact profitability and harm our business.

We face various risks and uncertainties as a result of our voluntary Titan XL recall program, including harm to our business, reputation, financial position and results of operations.

We discovered that under specific conditions, such as infrequent use or operators not following the prescribed maintenance program outlined in the owner's manual, certain component parts of one of our hand pieces — the Titan XL — could become defective. As a result, the defective hand piece could produce energy above that stated on the device setting, which could cause injury to patients, including redness, erythema, blisters and burns. In response to discovering this issue, we initiated a voluntary recall of our global installed base of Titan XL hand pieces starting May 2010 and have now completed the recall. Following a voluntary recall, there is generally an increase in product liability lawsuits filed against the Company.

If there were patients injured due to the malfunctioning of one of our voluntarily recalled Titan XL hand pieces that we are not presently aware of, we could incur significant legal fees and expenses in defending and/or settling such claims. Although we maintain product liability insurance and accrue for the cost of our deductible under such policies, there is no assurance that the entire loss of known claims or claims not known as of this point will be covered by our insurance. Each potential claim is evaluated on a case by case basis. At December 31, 2010, we have accrued for the cost of our deductible under our product liability insurance policy for all known claims. However, we have not recognized any potential contingent liability that may exceed our insurance coverage associated with any past or future lawsuits that may arise, as this is not estimable.

Our Titan hand piece refill revenue decreased 31% in 2010, compared to 2009. This decrease was due primarily to our voluntary recall of our Titan XL hand piece in 2010, in which we provided our eligible customers with a fully refilled Titan XL hand piece. If customers do not return to using their Titan hand pieces due to this voluntary recall, our Titan refill revenue could be negatively impacted in the future.

If either of the above mentioned two risks materializes, our revenue and profitability may be adversely affected, could materially harm our business, and our stock price could decline.

Federal regulatory reforms and changes occurring at the U.S. Food and Drug Administration, or FDA, could adversely affect our ability to sell our products profitably and financial condition.

From time to time, legislation is drafted and introduced in Congress that could significantly change the statutory provisions governing the clearance or approval, manufacture and marketing of a device. It is impossible to predict whether legislative changes will be enacted or FDA regulations, guidance or interpretations changed, and what the impact of such changes, if any, may be.

In addition, FDA regulations and guidance are often revised or reinterpreted by the agency in ways that may significantly affect our business and our products. Changes in FDA regulations may lengthen the regulatory approval process for medical devices and require additional clinical data to support regulatory clearance for the sale and marketing of our new products. In addition, it may require additional safety monitoring, labeling changes, restrictions on product distribution or use, or other measures after the introduction of our products to market. Either of these changes lengthen the duration to market, increase our costs of doing business, adversely affect the future permitted uses of approved products, or otherwise adversely affect the market for our products.

If we fail to obtain or maintain necessary FDA clearances for our products and indications, if clearances for future products and indications are delayed or not issued, if there are federal or state level regulatory changes or if we are found to have violated applicable FDA marketing rules, our commercial operations would be harmed.

Our products are medical devices that are subject to extensive regulation in the United States by the FDA for manufacturing, labeling, sale, promotion, distribution and shipping. Before a new medical device, or a new use of or labeling claim for an existing product, can be marketed in the United States, it must first receive either 510(k) clearance or pre-marketing approval from the FDA, unless an exemption applies. Either process can be expensive and lengthy. In the event that we do not obtain FDA clearances or approvals for our products, our ability to market and sell them in the United States and revenue derived there from may be adversely affected.

Medical devices may be marketed in the United States only for the indications for which they are approved or cleared by the FDA. For example, our recently introduced GenesisPlus product has a number of general indications for use in the U.S. that allows us to market the product in the U.S., however we can only market it internationally for the treatment of toenail fungus as it has a CE Mark approval. Another example is our Pearl Fractional product which is cleared only for skin resurfacing in the U.S. and our Titan product only for deep heating for the temporary relief of muscle aches and pains in the U.S. Therefore, we are prevented from promoting or advertising GenesisPlus, Titan and Pearl Fractional in the United States for any other indications. If we fail to comply with these regulations, it could result in enforcement action by the FDA which could lead to such consequences as warning letters, adverse publicity, criminal enforcement action and/or third-party civil litigation, each of which could adversely affect us.

We have obtained 510(k) clearance for the indications for which we market our products. However, our clearances can be revoked if safety or effectiveness problems develop. We also are subject to Medical Device Reporting regulations, which require us to report to the FDA if our products cause or contribute to a death or serious injury, or malfunction in a way that would likely cause or contribute to a death or serious injury. Our products are also subject to state regulations, which are, in many instances frequently changing. Changes in state regulations may impede sales. For example, federal regulations allow our products to be sold to, or on the order of, "licensed practitioners," as determined on a state-by-state basis. As a result, in some states, non-physicians may legally purchase our products. However, a state could change its regulations at any time, thereby disallowing sales to particular types of end users. We cannot predict the impact or effect of future legislation or regulations at the federal or state levels.

The FDA and state authorities have broad enforcement powers. If we fail to comply with applicable regulatory requirements, it could result in enforcement action by the FDA or state agencies, which may include any of the following sanctions:

- Warning letters, fines, injunctions, consent decrees and civil penalties;
- Repair, replacement, recall or seizure of our products;
- Operating restrictions or partial suspension or total shutdown of production;
- Refusing our requests for 510(k) clearance or pre-market approval of new products, new intended uses, or modifications to existing products;
- Withdrawing 510(k) clearance or pre-market approvals that have already been granted; and
- Criminal prosecution.

If any of these events were to occur, it could harm our business.

If we fail to comply with the FDA's Quality System Regulation and laser performance standards, our manufacturing operations could be halted, and our business would suffer.

We are currently required to demonstrate and maintain compliance with the FDA's Quality System Regulation, or QSR. The QSR is a complex regulatory scheme that covers the methods and documentation of the design, testing, control, manufacturing, labeling, quality assurance, packaging, storage and shipping of our products. Because our products involve the use of lasers, our products also are covered by a performance standard for lasers set forth in FDA regulations. The laser performance standard imposes specific record-keeping, reporting, product testing and product labeling requirements. These requirements include affixing warning labels to laser products, as well as incorporating certain safety features in the design of laser products. The FDA enforces the QSR and laser performance standards through periodic unannounced inspections. We had a full quality system audit in 2008 and an FDA audit of compliance with laser performance standards in 2010. There were no significant findings as a result of these audits and our responses have been accepted by the FDA. Our failure to take satisfactory corrective action in response to an adverse QSR inspection or our failure to comply with applicable laser performance standards could result in enforcement actions, including a public warning letter, a shutdown of our manufacturing operations, a recall of our products, civil or criminal penalties, or other sanctions, such as those described in the preceding paragraph, which would cause our sales and business to suffer.

If we modify one of our FDA-approved devices, we may need to seek re-approval, which, if not granted, would prevent us from selling our modified products or cause us to redesign our products.

Any modifications to an FDA-cleared device that would significantly affect its safety or effectiveness or that would constitute a major change in its intended use would require a new 510(k) clearance or possibly a pre-market approval. We may not be able to obtain additional 510(k) clearance or pre-market approvals for new products or for modifications to, or additional indications for, our existing products in a timely fashion, or at all. Delays in obtaining future clearance would adversely affect our ability to introduce new or enhanced products in a timely manner, which in turn would harm our revenue and future profitability.

We have made modifications to our devices in the past and may make additional modifications in the future that we believe do not or will not require additional clearance or approvals. If the FDA disagrees, and requires new clearances or approvals for the modifications, we may be required to recall and to stop marketing the modified devices, which could harm our operating results and require us to redesign our products.

We may be unable to obtain or maintain international regulatory qualifications or approvals for our current or future products and indications, which could harm our business.

Sales of our products outside the United States are subject to foreign regulatory requirements that vary widely from country to country. In addition, exports of medical devices from the United States are regulated by the FDA. Complying with international regulatory requirements can be an expensive and time-consuming process and approval is not certain. The time required for obtaining clearance or approvals, if required by other countries, may be longer than that required for FDA clearance or approvals, and requirements for such clearances or approvals may significantly differ from FDA requirements. We may be unable to obtain or maintain regulatory qualifications, clearances or approvals in other countries. We may also incur significant costs in attempting to obtain and in maintaining foreign regulatory approvals or qualifications. If we experience delays in receiving necessary qualifications, clearances or approvals to market our products outside the United States, or if we fail to receive those qualifications, clearances or approvals, we may be unable to market our products or enhancements in international markets effectively, or at all, which could have a material adverse effect on our business and growth strategy.

Product liability suits could be brought against us due to a defective design, material or workmanship or misuse of our products and could result in expensive and time-consuming litigation, payment of substantial damages and an increase in our insurance rates.

If our products are defectively designed, manufactured or labeled, contain defective components or are misused, we may become subject to substantial and costly litigation by our customers or their patients. Misusing our products or failing to adhere to operating guidelines could cause significant eye and skin damage, and underlying tissue damage. In addition, if our operating guidelines are found to be inadequate, we may be subject to liability. We have been involved, and may in the future be involved, in litigation related to the use of our products. Product liability claims could divert management's attention from our core business, be expensive to defend and result in sizable damage awards against us. We may not have sufficient insurance coverage for all future claims. We may not be able to obtain insurance in amounts or scope sufficient to provide us with adequate coverage against all potential liabilities. Any product liability claims brought against us, with or without merit, could increase our product liability insurance rates or prevent us from securing continuing coverage, could harm our reputation in the industry and could reduce product sales. In addition, we historically experienced steep increases in our product liability insurance premiums as a percentage of revenue. If our premiums continue to rise, we may no longer be able to afford adequate insurance coverage.

If customers are not trained and / or our products are used by non-physicians, it could result in product misuse and adverse treatment outcomes, which could harm our reputation, result in product liability litigation, distract management, result in additional costs, all of which could harm our business.

Because we do not require training for users of our products, and sell our products at times to non-physicians, there exists an increased potential for misuse of our products, which could harm our reputation and our business. U.S. federal regulations allow us to sell our products to or on the order of "licensed practitioners." The definition of "licensed practitioners" varies from state to state. As a result, our products may be purchased or operated by physicians with varying levels of training, and in many states, by non-physicians, including nurse practitioners, chiropractors and technicians. Outside the United States, many jurisdictions do not require specific qualifications or training for purchasers or operators of our products. We do not supervise the procedures performed with our products, nor do we require that direct medical supervision occur. We and our distributors generally offer but do not require product training to the purchasers or operators of our products. In addition, we sometimes sell our systems to companies that rent our systems to third parties and that provide a technician to perform the procedures. The lack of training and the purchase and use of our products by non-physicians may result in product misuse and adverse treatment outcomes, which could harm our reputation and our business, and, in the event these result in product liability litigation, distract management and subject us to liability, including legal expenses.

We have recently entered into strategic alliances to distribute third party products internationally. To successfully market and sell these products, we must address many issues that are unique to these businesses and could reduce our available cash reserves and negatively impacting our profitability.

Recently, we have entered into distribution arrangements pursuant to which we utilize our sales force and distributors to sell products manufactured by other companies. We entered into an agreement with Obagi to distribute certain of their proprietary cosmeceuticals, or skin care products, in Japan. This agreement requires us to purchase an annual minimum dollar amount of their product. During 2010 the initial year of the agreement, the minimum purchase requirement was \$1.25 million. The minimum purchase requirement for 2011 and beyond has yet to be finalized but is expected to be consistent with 2010. If we do not make these minimum purchases, we could lose exclusivity for distributing Obagi products to physicians in Japan. In addition, in December 2009, we entered into an agreement with Sound Surgical Technologies, Inc. to distribute their VASER® Lipo System in certain European countries and Canada. However, in September 2010, we decided to discontinue distributing the VASER product. Finally, we also have an agreement with BioForm Medical Inc. (acquired by Merz Pharma Group in January 2009), to distribute their Radiesse® dermal filler product in Japan. Each of these distribution agreements presents its own unique risks and challenges. For example, to sell products in partnership with Obagi we need to invest in creating a sales structure that is experienced in the sale of cosmeceuticals and not in capital equipment. We need to commit resources to training this sales force, obtaining regulatory licenses in Japan and developing new marketing materials to promote the sale of Obagi products. For each of these distribution arrangements, until we can develop our own experienced sales force, we may need to pay third party distributors to sell the products which will result in higher fees and lower margins than if we sell direct to customers. In addition, the minimum commitments and other costs of distributing products manufactured by these companies may exceed the incremental revenue that we derive from the sale of their products thereby reducing our available cash reserves and negatively impacting our profitability.

To successfully market and sell our products internationally, we must address many issues that are unique to our international business.

Our international revenue was \$33.9 million or 64% of our total revenue in 2010. International revenue is a material component of our business strategy. We depend on third-party distributors and a direct sales force to sell our products internationally, and if they underperform, we may be unable to increase or maintain our level of international revenue. To grow our business, we will need to improve productivity in current sales territories and expand into new territories. However, direct sales productivity may not improve and distributors may not accept our business or commit the necessary resources to market and sell our products to the level of our expectations. As a result, we may not be able to increase or maintain international revenue growth.

We believe, as we continue to manage our international operations and develop opportunities in additional international territories, our international revenue will be subject to a number of risks, including:

- Difficulties in staffing and managing our foreign operations;
- Export restrictions, trade regulations and foreign tax laws;
- Fluctuating foreign currency exchange rates;
- Foreign certification and regulatory requirements;
- Lengthy payment cycles and difficulty in collecting accounts receivable;
- Customs clearance and shipping delays;
- Political and economic instability;
- Lack of awareness of our brand in international markets;
- Preference for locally-produced products; and
- Reduced protection for intellectual property rights in some countries.

If one or more of these risks were realized, it could require us to dedicate significant resources to remedy the situation; and if we were unsuccessful at finding a solution, we may not be able to sell our products in a particular market and, as a result, our revenue may decline.

If PSS World Medical fails to perform to our expectations, we may fail to achieve anticipated operating results.

We have a distribution agreement with PSS World Medical. PSS sales professionals work in coordination with our sales force to locate new customers for our products throughout the United States. Revenue from PSS has significantly declined since 2008. Our revenue from PSS, as a percentage of worldwide revenue, was 5% in 2010, 7% in 2009 and 14% in 2008. Although we continue to work closely with, and focus our attention on, our PSS relationship, there is no assurance that this will translate into increased revenue for us. Further, if revenue from PSS does not improve, or if they terminate our relationship, it may have a significant adverse effect on our revenue, financial condition and results of operations.

Adverse conditions in the global banking industry and credit markets may adversely impact the value of our marketable investments or impair our liquidity.

We invest our excess cash primarily in money market funds and in highly liquid debt instruments of the U.S. government and its agencies, and U.S. municipalities. As of December 31, 2010, our balance in marketable investment was \$77.5 million. The longer the duration of a security, the more susceptible it is to changes in market interest rates and bond yields. As yields increase, those securities with a lower yield-at-cost show a mark-to-market unrealized loss. For example, assuming a hypothetical increase in interest rates of one percentage point, the fair value of our total investment portfolio as of December 31, 2010 would have potentially decreased by approximately \$651,000, resulting in an unrealized loss that would subsequently adversely impact our earnings. As a result, changes in the market interest rates will affect our future net income (loss).

We may be required to record impairment charges in future quarters as a result of the decline in value of our longterm investments in auction rate securities (ARS).

Included under the caption of "Long-term investments" in the Consolidated Balance Sheet as of December 31, 2010 are \$6.8 million of ARS. These ARS were designed to provide liquidity through an auction process that resets the applicable interest rate at predetermined calendar intervals, generally every 35 days. Though approximately \$5.1 million (par value) of our original holdings of \$13.4 million (par value) of ARS have been redeemed at full par value since 2008, auctions for the remaining ARS in our portfolio at December 31, 2010 continue to fail and they remain as illiquid. Upon an auction failure, the interest rates do not reset at a market rate but instead reset based on a formula contained in the prospectus of the individual security, which rate is generally higher than the prevailing market rate. The failure of the auctions impacts our ability to readily liquidate our ARS into cash until a future auction of these investments is successful, a buyer is found outside of the auction process, or the ARS is refinanced by the issuer into another type of debt instrument.

If there is a decline in fair value in our ARS that is considered other-than-temporary then we would have to record an impairment charge in our Consolidated Statement of Operations for the loss in value associated with the worsening of the credit worthiness (credit losses) of the issuer, which would reduce future earnings, harm our business and may cause our stock price to decline.

We depend on skilled and experienced personnel to operate our business effectively. If we are unable to recruit, hire, train and retain these employees, our ability to manage and expand our business will be harmed, which would impair our future revenue and profitability.

Our success largely depends on the skills, experience and efforts of our officers and other key employees. Except for Change of Control and Severance Agreements for our executive officers, we do not have employment contracts with any of our officers or other key employees. Any of our officers and other key employees may terminate their employment at any time. We do not have a succession plan in place for each of our officers and key employees. In addition, we do not maintain "key person" life insurance policies covering any of our employees. The loss of any of our senior management team members could weaken our management expertise and harm our business.

Our ability to retain our skilled labor force and our success in attracting and hiring new skilled employees are critical factors in determining whether we will be successful in the future. We may not be able to meet our future hiring needs or retain existing personnel. We may face particularly significant challenges and risks in hiring, training, managing and retaining engineering and sales and marketing employees. Failure to attract, train and retain personnel, particularly technical and sales and marketing personnel, would materially harm our ability to compete effectively and grow our business.

The price of our common stock may fluctuate substantially. We have a limited number of shares of common stock outstanding, a large portion of which is held by a small number of investors, which could result in the increase in volatility of our stock price.

As of December 31, 2010, approximately 51% of our outstanding shares of common stock were held by 10 institutional investors. As a result of our relatively small public float, our common stock may be less liquid than the stock of companies with broader public ownership. Among other things, trading of a relatively small volume of our common stock may have a greater impact on the trading price for our shares than would be the case if our public float were larger.

The public market price of our common stock has in the past fluctuated substantially and, due to the current concentration of stockholders, it may continue to do so in the future. The market price for our common stock could also be affected by a number of other factors, including:

- The general market conditions unrelated to our operating performance;
- Sales of large blocks of our common stock, including sales by our executive officers, directors and our large institutional investors;
- Quarterly variations in our, or our competitors', results of operations;
- Changes in analysts' estimates, investors' perceptions, recommendations by securities analysts or our failure to achieve analysts' estimates;
- The announcement of new products or service enhancements by us or our competitors;
- The announcement of the departure of a key employee or executive officer by us or our competitor;
- Regulatory developments or delays concerning our, or our competitors' products; and
- The initiation of litigation by us or against us.

Actual or perceived instability in our stock price could reduce demand from potential buyers of our stock, thereby causing our stock price to either remain depressed or to decline further.

We may be involved in future costly intellectual property litigation, which could impact our future business and financial performance,

Our competitors or other patent holders may assert that our present or future products and the methods we employ are covered by their patents. In addition, we do not know whether our competitors own or will obtain patents that they may claim prevent, limit or interfere with our ability to make, use, sell or import our products. Although we may seek to resolve any potential future claims or actions, we may not be able to do so on reasonable terms, or at all. If, following a successful third-party action for infringement, we cannot obtain a license or redesign our products, we may have to stop manufacturing and selling the applicable products and our business would suffer as a result. In addition, a court could require us to pay substantial damages, and prohibit us from using technologies essential to our products, any of which would have a material adverse effect on our business, results of operations and financial condition.

We may become involved in litigation not only as a result of alleged infringement of a third party's intellectual property rights but also to protect our own intellectual property. For example, we have been, and may hereafter become, involved in litigation to protect the trademark rights associated with our company name or the names of our products. Infringement and other intellectual property claims, with or without merit, can be expensive and time-consuming to litigate, and could divert management's attention from our core business.

Any acquisitions that we make could disrupt our business and harm our financial condition.

From time to time we evaluate potential strategic acquisitions of complementary businesses, products or technologies. We may also consider joint ventures and other collaborative projects. We may not be able to identify appropriate acquisition candidates or strategic partners, or successfully negotiate, finance or integrate any businesses, products or technologies that we acquire. Furthermore, the integration of any acquisition and management of any collaborative project may divert management's time and resources from our core business and disrupt our operations and we may incur significant legal, accounting and banking fees in connection with such a transaction. In addition, if we purchase a company that is not profitable, our cash balances may be reduced or depleted. We do not have any experience as a team with acquiring companies or products. If we decide to expand our product offerings beyond laser and light-based products, we may spend time and money on projects that do not increase our revenue. Any cash acquisition we pursue would diminish our available cash balances to us for other uses, and any stock acquisition could be dilutive to our stockholders.

While we from time to time evaluate potential acquisitions of businesses, products and technologies, and anticipate continuing to make these evaluations, we have no present understandings, commitments or agreements with respect to any material acquisitions or collaborative projects.

Our manufacturing operations are dependent upon third-party suppliers, making us vulnerable to supply shortages and price fluctuations, which could harm our business.

Many of the components and materials that comprise our products are currently manufactured by a limited number of suppliers. A supply interruption or an increase in demand beyond our current suppliers' capabilities could harm our ability to manufacture our products until a new source of supply is identified and qualified. Our reliance on these suppliers subjects us to a number of risks that could harm our business, including:

- Interruption of supply resulting from modifications to or discontinuation of a supplier's operations;
- Delays in product shipments resulting from uncorrected defects, reliability issues or a supplier's variation in a component;
- A lack of long term supply arrangements for key components with our suppliers;
- Inability to obtain adequate supply in a timely manner, or on reasonable terms;
- Difficulty locating and qualifying alternative suppliers for our components in a timely manner;
- Production delays related to the evaluation and testing of products from alternative suppliers and corresponding regulatory qualifications; and
- Delay in supplier deliveries.

Any interruption in the supply of components or materials, or our inability to obtain substitute components or materials from alternate sources at acceptable prices in a timely manner, could impair our ability to meet the demand of our customers, which would have an adverse effect on our business.

Intellectual property rights may not provide adequate protection for some or all of our products, which may permit third parties to compete against us more effectively.

We rely on patent, copyright, trade secret and trademark laws and confidentiality agreements to protect our technology and products. At December 31, 2010, we had 17 issued U.S. patents. Some of our components, such as our laser module, electronic control system and high-voltage electronics, are not, and in the future may not be, protected by patents. Additionally, our patent applications may not issue as patents or, if issued, may not issue in a form that will be advantageous to us. Any patents we obtain may be challenged, invalidated or legally circumvented by third parties. Consequently, competitors could market products and use manufacturing processes that are substantially similar to, or superior to, ours. We may not be able to prevent the unauthorized disclosure or use of our technical knowledge or other trade secrets by consultants, vendors, former employees or current employees, despite the existence generally of confidentiality agreements and other contractual restrictions. Monitoring unauthorized uses and disclosures of our intellectual property is difficult, and we do not know whether the steps we have taken to protect our intellectual property rights to the same extent as the laws of the United States.

The absence of complete intellectual property protection exposes us to a greater risk of direct competition. Competitors could purchase one of our products and attempt to replicate some or all of the competitive advantages we derive from our development efforts, design around our protected technology, or develop their own competitive technologies that fall outside of our intellectual property rights. If our intellectual property is not adequately protected against competitors' products and methods, our competitive position and our business could be adversely affected.

Healthcare reform legislation could adversely affect our future profitability and financial condition.

The President and members of Congress passed legislation relating to healthcare reform. Our products are not reimbursed by insurance companies or federal or state governments and some of this legislation will, therefore, not affect us. This legislation, however, does include several aspects that will apply to us, including a tax on our U.S. revenue which is applicable to us beginning in 2013. While we are presently evaluating the full scope of how this legislation will impact our operations, including how to administer this tax, we believe this will adversely affect our future profitability and financial condition.

We offer credit terms to some qualified customers and also to leasing companies to finance the purchase of our products. In the event that any of these customers default on the amounts payable to us, our earnings may be adversely affected.

While we qualify customers to whom we offer credit terms (generally net 30 to 60 days), we cannot provide any assurance that the financial position of these customers will not change adversely before we receive payment. Our general and administrative expenses and earnings are negatively impacted by customer defaults and cause an increase in the allowance for doubtful accounts. In the event that there is a default by any customers to whom we have provided credit terms in the future, we may recognize a bad debt charge in our general and administrative expenses and this could negatively affect our earnings and results of operations.

We are subject to fluctuations in the exchange rate of the U.S. dollar and foreign currencies.

As a result of recent fluctuations in currency markets and the strong dollar relative to many other major currencies, our products priced in U.S. dollars may be more expensive relative to products of our foreign competitors, which could result in lower revenue. We do not actively hedge our exposure to currency rate fluctuations. While we transact business primarily in U.S. Dollars, and a significant proportion of our revenue is denominated in U.S. Dollars, a portion of our costs and revenue is denominated in other currencies, such as the Euro, Japanese Yen, Australian Dollar, Canadian Dollar and British Pound Sterling. As a result, changes in the exchange rates of these currencies to the U.S. Dollar will affect our net income (loss).

The expense and potential unavailability of insurance coverage for our customers could adversely affect our ability to sell our products, and therefore our financial condition.

Some of our customers and prospective customers have had difficulty in procuring or maintaining liability insurance to cover their operation and use of our products. Medical malpractice carriers are withdrawing coverage in certain states or substantially increasing premiums. If this trend continues or worsens, our customers may discontinue using our products and potential customers may opt against purchasing laser and light based products due to the cost or inability to procure insurance coverage. The unavailability of insurance coverage for our customers and prospects could adversely affect our ability to sell our products, and that could harm our financial condition.

Anti-takeover provisions in our Amended and Restated Certificate of Incorporation and Bylaws, and Delaware law, contain provisions that could discourage a takeover.

Our Amended and Restated Certificate of Incorporation and Bylaws, and Delaware law, contain provisions that might enable our management to resist a takeover, and might make it more difficult for an investor to acquire a substantial block of our common stock. These provisions include:

- A classified board of directors:
- Advance notice requirements to stockholders for matters to be brought at stockholder meetings;
- A supermajority stockholder vote requirement for amending certain provisions of our Amended and Restated Certificate of Incorporation and bylaws;

- Limitations on stockholder actions by written consent; and
- The right to issue preferred stock without stockholder approval, which could be used to dilute the stock ownership of a potential hostile acquirer.

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These provisions, as well as Change of Control and Severance Agreements entered into with each of our executive officers, might discourage, delay or prevent a change in control of our company or a change in our management. The existence of these provisions could adversely affect the voting power of holders of common stock and limit the price that investors might be willing to pay in the future for shares of our common stock.

ITEM 1B. UNRESOLVED STAFF COMMENTS

Not applicable.

ITEM 2. PROPERTIES

Our corporate headquarters and U.S. operations are located in an approximately 66,000 square foot facility in Brisbane, California. We lease these premises under a non-cancelable operating lease which expires on December 31, 2017. In addition, we have leased office facilities in certain international countries as follows:

Country	Square Footage	Lease termination or Expiration
Japan	Approximately 5,790	Three leases of which two expire in May 2012 and one expires in
•		July 2013, but may be cancelled at any time with a six-month notice.
Switzerland	Approximately 3,174	One lease which expires in March 2013.
France	Approximately 450	Lease expires in November 2011, but may be cancelled at any time with a three-month notice.
Spain	Approximately 175	Lease automatically renews at the end of each six-month period.

We believe that these facilities are adequate for our current and future needs for at least the next twelve months.

ITEM 3. LEGAL PROCEEDINGS

A Telephone Consumer Protection Act, or TCPA, class action lawsuit was filed against us in January 2008 and settled in 2009 on a class-wide basis. We paid a total of \$950,000 in exchange for a full release of all claims and recorded a net charge of \$850,000 in our 2009 Consolidated Statements of Operations for the cost of the settlement, net of the administrative expenses and contributions from our insurance carrier. All monies were distributed in conjunction with this settlement, a final accounting hearing was presented to the Court on February 3, 2011, and the case closed.

Two securities class action lawsuits were filed against us and two of our executive officers in 2007. The plaintiffs claimed to represent purchasers of our common stock from January 31, 2007 through May 7, 2007 and generally alleged that materially false statements and omissions were made regarding our financial prospects. In 2008, the Court issued an order dismissing the plaintiffs' consolidated, amended complaint without prejudice and closed the case on its own initiative. On November 26, 2008, the plaintiffs filed a Notice of Appeal to the U.S. Court of Appeals for the Ninth Circuit. In 2009, the parties filed their written briefs with that Court. In 2010, both parties presented oral arguments and the Court affirmed the District Court's order dismissing the complaint.

ITEM 4. [RESERVED]

PART II

ITEM 5. MARKET FOR THE REGISTRANT'S COMMON EQUITY, RELATED STOCKHOLDER MATTERS AND ISSUER PURCHASES OF EQUITY SECURITIES

Stock Exchange Listing

Our common stock trades on The NASDAQ Global Select Market under the symbol "CUTR." As of February 28, 2011, the closing sale price of our common stock was \$9.66 per share.

Common Stockholders

We had 11 stockholders of record as of February 28, 2011. Since many stockholders choose to hold their shares under the name of their brokerage firm, we believe, the actual number of stockholders was in excess of 2600.

Stock Prices

The following table sets forth quarterly high and low closing sales prices of our common stock for the indicated fiscal periods:

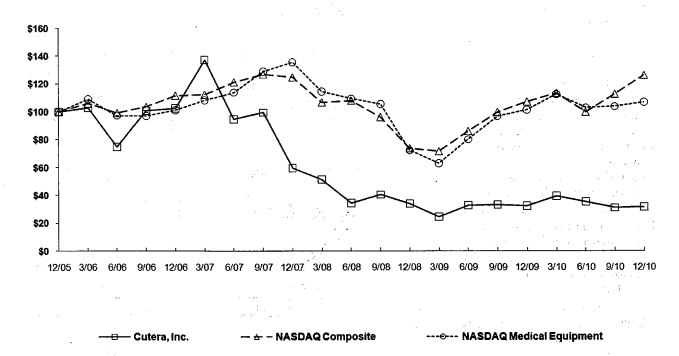
		Common Stock										
		20	10			20	09					
		ligh		Low	F	Iigh		Low				
4th Quarter	\$	8.39	\$	7.01	\$	9.63	\$	7.97				
3rd Quarter		9,00		6.99		9.40	•	7.85				
2nd Quarter	·	12.04		8.62		9.03		5.93				
1st Quarter		11.03		8.25		8.71		5.57				

Performance Graph

Below is a graph showing the cumulative total return to our stockholders during the period from December 31, 2005 through December 31, 2010 in comparison to the cumulative return on the NASDAQ Composite Index (U.S.) and the NASDAQ Medical Equipment Index during that same period. (1) The results assume that \$100 was invested on December 31, 2005.

COMPARISON OF 5 YEAR CUMULATIVE TOTAL RETURN*

Among Cutera, Inc., the NASDAQ Composite Index and the NASDAQ Medical Equipment Index



*\$100 invested on 12/31/05 in stock or index, including reinvestment of dividends. Fiscal year ending December 31.

The information under "Performance Graph" is not deemed filed with the Securities and Exchange Commission and is not to be incorporated by reference in any filing of Cutera under the Securities Act of 1933, as amended, or the Securities Exchange Act of 1934, as amended, whether made before or after the date of this 10-K and irrespective of any general incorporation language in those filings.

Dividend Policy

We have never paid a cash dividend and have no present intention to pay cash dividends in the foreseeable future. We intend to retain any future earnings for use in our business.

We did not sell any unregistered securities during the period covered by this Annual Report on Form 10-K.

The information required by this Item regarding equity compensation plans is incorporated by reference to the information set forth in Part III Item 12 of this Annual Report on Form 10-K.

Securities Authorized for Issuance under Equity Compensation Plans

See Part III, Item 12 for information regarding securities authorized for issuance under equity compensation plans.

ITEM 6. SELECTED FINANCIAL DATA

The table set forth below contains certain consolidated financial data for each of our last five fiscal years. This data should be read in conjunction with the detailed information, financial statements and related notes, as well as Management's Discussion and Analysis of Financial Condition and Results of Operations included elsewhere herein.

	Year Ended December 31,									
Consolidated Statements of Operations Data (in thousands, except per share data):		2010		2009		2000		400		••••
Net revenue	\$	53,274	\$	53,682	\$	2008 83,379	æ	2007 101,726	Ф	2006 100,692
Cost of revenue.	Ψ	23,058	Ψ	21,759		32,358	Ф	35,002	Φ	•
Gross profit	_	30,216	_	31,923	_	51,021	_	66,724	_	29,859 70,833
Operating expenses:	_	30,210	_	31,923	_		_	00,724	_	70,833
Sales and marketing		24,735		24,286		35,354		38,277		32,890
Research and development		7,004		6,810		7,550		7,169		52,890 6,473
General and administrative		9,576		10,320		11,270		11,721		15,192
Litigation settlement		9,570		850		11,2/0		11,/21		18,935
Total operating expenses	_	41,315	_	42,266		54,174	_	57,167	_	73,490
Income (loss) from operations		(11,099)	_	$\frac{42,200}{(10,343)}$	_	(3,153)	_	9,557	_	
Interest and other income, net.		583		1,572		3,046		4,207		(2,657)
Other-than-temporary impairments of long-term		363		1,372		3,040		4,207		3,596
investments						(3,554)				
Income (loss) before income taxes	_	(10,516)	-	(8,771)	_	(3,661)	_	13,764		939
Provision (benefit) for income taxes		2		8,908		(3,001)		3,260		
Net income (loss)	\$	(10,518)	\$	(17,679)	\$	(2,869)	\$		<u>~</u>	(1,184)
Net income (loss) available to common	. <u>Ф</u>	(10,516)	<u> </u>	(17,079)	Ф	(2,809)	<u> </u>	10,504	\$	2,123
stockholders used in basic net income per	•	•								
share	•	(10.510)	Φ	(17 (70)	ø	(2.9(0)	Ø	10.504	Φ	0.100
	<u>\$</u>	(10,518)	<u>\$</u>	<u>(17,679</u>)	<u>\$</u>	(2,869)	\$	10,504	<u>\$</u>	2,123
Net income (loss) per share:	Ф	(0.70)	Φ	(1.22)	Φ.	(0.00)	•		•	0.4=
Basic	\$	(0.78)	\$	(1.33)	\$	(0.22)	\$	0.80	\$	0.17
Diluted	\$	(0.78)	\$	(1.33)	\$	(0.22)	\$	0.74	\$	0.15
Weighted-average number of shares used in per		. 1								
share calculations:	•			* **				٠.		
Basic	_	13,540	_	13,279		12,770	_	13,153		12,558
Diluted	1 11	13,540		13,279		12,770		14,228	: .	14,278
										
Constituted Balance Charles and the Constitution of the Constituti		2010			s of	December 3	1,			
Consolidated Balance Sheet Data (in thousands): Cash and cash equivalents	\$	2010 7	_	2009 -	<u>.</u>	2008	<u>.</u>	2007		2006
Marketable investments	•	12,519				36,540		11,054	\$	11,800
		77,484		,	٠,	60,653		88,510		96,285
Long-term investments Working capital (current assets less current		6,784		7,275		9,627		7,429	ź.	#**** **
liabilities)		90,339		06.015		101 (44		106 904		111.000
Total assets		111,805		96,015 121,352		101,644		106,894		111,999
Retained earnings.		6,736		,		137,476		138,653		133,875
Total stockholders' equity		,		17,254		31,410		34,279		23,866
Total Stockholders Equity		95,417		100,853		112,108		109,353	- '	109,732

ITEM 7. MANAGEMENT'S DISCUSSION AND ANALYSIS OF FINANCIAL CONDITION AND RESULTS OF OPERATIONS

The following discussion should be read in conjunction with our audited financial statements and notes thereto for the fiscal year ended December 31, 2010. This Annual Report on Form 10-K, including the following sections, contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Throughout this Report, and particularly in this Item 7, the forward-looking statements are based upon our current expectations, estimates and projections and that reflect our beliefs and assumptions based upon information available to us at the date of this Report. In some cases, you can identify these statements by words such as "may," "might," "will," "should," "expects," "plans," "anticipates," "believes," "estimates," "predicts," "potential" or "continue," and other similar terms. These forward-looking statements are not guarantees of future performance and are subject to risks, uncertainties, and assumptions that are difficult to predict. Our actual results, performance or achievements could differ materially from those expressed or implied by the forward-looking statements. The forward-looking statements include, but are not limited to, statements relating to our future financial performance, the ability to grow our business, increase our revenue, manage expenses, generate additional cash, achieve and maintain profitability, develop and commercialize existing and new products and applications, improve the performance of our worldwide sales and distribution network, and to the outlook regarding long term prospects. We caution you not to place undue reliance on these forward-looking statements, which reflect management's analysis only as of the date of this Annual Report on Form 10-K. We undertake no obligation to update forward-looking statements to reflect events or circumstances occurring after the date of this Form 10-K.

Some of the important factors that could cause our results to differ materially from those in our forward-looking statements, and a discussion of other risks and uncertainties, are discussed in Item 1A—Risk Factors commencing on page 13. We encourage you to read that section carefully as well as other risks detailed from time to time in our filings with the SEC.

Introduction

The Management's Discussion and Analysis, or MD&A, is organized as follows:

- Executive Summary. This section provides a general description and history of our business, a brief discussion of our product lines and the opportunities, trends, challenges and risks we focus on in the operation of our business.
- Critical Accounting Policies and Estimates. This section describes the key accounting policies that are affected by critical accounting estimates.
- Recent Accounting Guidance. This section describes the issuance and effect of new accounting pronouncements that are and may be applicable to us.
- Results of Operations. This section provides our analysis and outlook for the significant line items on our Consolidated Statements of Operations.
- Liquidity and Capital Resources. This section provides an analysis of our liquidity and cash flows, as well as a discussion of our commitments that existed as of December 31, 2010.

Executive Summary

Company Description. We are a global medical device company specializing in the design, development, manufacture, marketing and servicing of laser and light-based aesthetics systems for practitioners worldwide. We offer easy-to-use products based on five platforms — CoolGlide[®], Xeo[®], Solera[®], GenesisPlusTM and Excel VTM — each of which enables physicians and other qualified practitioners to perform safe and effective aesthetic procedures for their customers.

Our corporate headquarters and U.S. operations are located in Brisbane, California, from where we conduct our manufacturing, warehousing, research and development, regulatory, sales and marketing, service, and administrative activities. In the United States, we market, sell and service our products through direct sales and service employees, and a distribution relationship with PSS World Medical Shared Services, Inc. ("PSS"), a wholly owned subsidiary of PSS World Medical which has over 700 sales representatives serving physician offices throughout the United States. We also sell certain items such as our Titan hand piece refills and marketing brochures online over the internet.

International sales are generally made through direct sales employees and a worldwide distributor network in over 35 countries. Outside of the United States, we have a direct sales presence in Australia, Canada, France, Japan, Spain, Switzerland and the United Kingdom.

Products. Our revenue is derived from the sale of Products, Upgrades, Service, Titan hand piece refills, and Dermal fillers and cosmeceutical products. Product revenue represents the sale of a system. A system consists of a console that incorporates a universal graphic user interface, a laser and/or light-based module, control system software and high voltage electronics; as well as one or more hand pieces. However, depending on the application, the laser or light-based module is sometimes contained in the hand piece such as with our Pearl and Pearl Fractional applications instead of within the console.

We offer our customers the ability to select the system that best fits their practice at the time of purchase and then to cost-effectively add applications to their system as their practice grows. This provides customers the flexibility to upgrade their systems whenever they want and provides us with a source of recurring revenue which we classify as Upgrade revenue. Service revenue relates to amortization of pre-paid service contract revenue and receipts for time and materials services on out-of-warranty products. Titan hand piece refill revenue is associated with our Titan hand piece which requires replacement of the optical source after a set number of pulses have been used. In Japan, we distribute BioForm, Inc.'s (BioForm) Radiesse® dermal filler product and Obagi's cosmeceutical products.

Significant Business Trends. We believe that our ability to grow revenue has been, and will continue to be, primarily dependent on the following:

- Continuing to expand our product offerings.
- Ongoing investment in our global sales and marketing infrastructure.
- Use of clinical results to support new aesthetic products and applications.
- Enhanced luminary development and reference selling efforts (to develop a location where our products can be displayed and used to assist in selling efforts).
- Customer demand for our products.
- Consumer demand for our products...
- Marketing to physicians in the core dermatology and plastic surgeon specialties, as well as outside those specialties.
- Generating ongoing revenue from our growing installed base of customers through the sale of Service,
 Upgrade, Titan hand piece refill, and Dermal fillers and cosmeceutical products.

Our U.S. revenue decreased by 8% and our international revenue increased by 4% in 2010, compared to 2009. International revenue as a percent of total revenue was 64% in 2010 and 61% in 2009. We believe the decline in U.S. revenues was attributable to several factors, including:

- Our Products and Upgrades ASPs in 2010 were lower than 2009 as a result of customers purchasing fewer applications for systems and lower pricing resulting from competitive discounting pressures.
- We experienced a temporary decline in our Titan refill revenue in 2010, compared to 2009, due to a voluntary recall of certain Titan XL hand pieces. All customers that had a Titan XL hand piece subject to the recall, were provided with fully refilled hand pieces, which delayed their purchase of a refill.

Our total international revenue increased by 4%, with growth being sourced from most of the international locations that we sell in. In Japan, our revenue increased by 41% in 2010, compared to 2009, as a result of growth from all of our product categories and the recently added Dermal fillers and cosmeceuticals business. With respect to Australia, our revenue declined in 2010 by approximately 65% due to 2009 being an unusually high revenue year in which a special tax incentive was offered by the government to incentivize customers to purchase capital equipment.

Our service revenue remained relatively unchanged in 2010, compared to 2009. Service contract amortization is a primary component of total service revenue. Our installed base of customers continued to increase in 2010 albeit at a slower pace with a bias toward international. Our deferred service revenue balance decreased by \$1.4 million, or 17%, to \$6.8 million as of December 31, 2010, compared to December 31, 2009. We believe the decline was attributable to:

• In years prior to 2009, we had discounted pricing promotions for incentivizing customers to purchase multiple year service contracts, however, in 2009 and 2010 we did not have such promotions.

- A decrease in unit sales volume in 2009 (but not in 2010), which historically included an element of deferred revenue for service contracts beyond our standard warranty terms.
- Recently, fewer customers have been purchasing extended service contracts in response to improved product reliability and the recent tougher economic environment.

Our gross margin in 2010 was 57%, compared with 59% in 2009 and 61% in 2008. This decline was due to several factors, including:

- The 2010 voluntary recall of certain Titan XL hand pieces whereby eligible customers were provided with fully refilled hand pieces;
- A \$1.7 million, or 31%, temporary decrease in our Titan refill revenue in 2010, compared to 2009, for which we traditionally earn a higher gross margin than our blended total gross margin percentage;
- Our ASPs have declined due primarily to customers purchasing fewer applications on their platforms and due to competitive discounting pressures; and
- A higher proportion of distributor revenue that carries a lower gross margin; partially offset by
- Lower manufacturing expenses resulting from headcount reductions; and
- Reduced warranty and service expenses as a result of improved product reliability (for products other than Titan XL hand pieces).

Our sales and marketing expenses remained relatively flat at \$24.7 million in 2010, compared with \$24.3 million in 2009. As a percentage of net revenue, in 2010 sales and marketing expenses were at 47%, compared to 45% in 2009. This slight increase was primarily attributable to the 8% lower revenue in the U.S. in 2010, compared to 2009.

Our research and development, or R&D, expenses increased slightly to \$7.0 million in 2010, compared with \$6.8 million in 2009. This increase was associated with higher personnel expenses resulting from higher headcount in engineering relating to new product development programs. As a percentage of net revenue, R&D expenses remained flat at 13% in 2010, compared to 2009.

Our general and administrative, or G&A, expenses decreased by approximately \$744,000, or 7%, to \$9.6 million in 2010, compared to 2009. As a percentage of net revenue, G&A expenses decreased slightly to 18% in 2010, compared to 19% in 2009. This decrease was due primarily to lower bad debt expenses, lower legal fees and litigation settlement costs.

In response to the economic conditions and a decline in revenue during 2009, we reduced our company-wide workforce by approximately 18% and implemented other cost-reduction measures in the first half of 2009. The headcount reductions impacted all departments and functions and resulted in restructuring charges of approximately \$880,000 in first half of 2009.

In 2009, we settled our TCPA class action lawsuit and in that regard recorded a charge of \$850,000 for the cost of the settlement, net of administrative expenses and amounts that were recovered from our insurance carrier. See "Item 3 - Legal Proceedings" in Part I, of this Form 10-K.

Factors that May Impact Future Performance

Our industry is impacted by numerous competitive, regulatory and other significant factors. Our industry is highly competitive and our future performance depends on our ability to compete successfully. Additionally, our future performance is dependent upon our ability to continue to expand our product offerings with innovative technologies, obtain regulatory clearances for our products, protect the proprietary technology of our products and our manufacturing processes, manufacture our products cost-effectively, and successfully market and distribute our products in a profitable manner. If we fail to execute on the aforementioned initiatives, our business would be adversely affected. A detailed discussion of these and other factors that could impact our future performance are provided in Part I, Item 1A "Risk Factors."

Critical Accounting Policies and Estimates

The preparation of our Consolidated Financial Statements and related disclosures in conformity with generally accepted accounting principles in the United States (GAAP) requires us to make estimates, judgments and assumptions that affect the reported amounts of assets, liabilities, revenue and expenses. These estimates, judgments and assumptions are based on historical experience and on various other factors that we believe are reasonable under the circumstances. We periodically review our estimates and make adjustments when facts and circumstances dictate. To the extent that there are material differences between these estimates and actual results, our financial condition or results of operations will be affected.

Critical accounting estimates, as defined by the Securities and Exchange Commission (SEC), are those that are most important to the portrayal of our financial condition and results of operations and require our management's most difficult and subjective judgments and estimates of matters that are inherently uncertain. Our critical accounting estimates are as follows:

Revenue Recognition

We recognize revenue from the sale of Products, Upgrades, Titan hand piece refills, and Dermal fillers and cosmeceuticals when title and risk of ownership has been transferred, provided that:

- Persuasive evidence of an arrangement exists;
- Delivery has occurred or services have been rendered;
- The fee is fixed or determinable; and
- Collectability is reasonably assured.

Determination of whether persuasive evidence of an arrangement exists and whether delivery has occurred or services have been rendered, are based on management's judgments regarding the fixed nature of the fee charged for services rendered and products delivered, and the collectability of those fees. In instances where final acceptance of the product is specified by the customer or collectability has not been reasonably assured, revenue is deferred until all acceptance criteria have been met. Revenue under service contracts is recognized on a straight-line basis over the period of the applicable service contract. Service revenue, not under a service contract, is recognized as the services are provided. Should changes in conditions cause management to determine these criteria are not met for certain future transactions, revenue recognized for any reporting period could be adversely affected.

Fair Value Measurement of our Long Term Auction Rate Securities Investments

We hold a variety of interest bearing auction rate securities (ARS) that represent investments in pools of student loan assets. At the time of acquisition, these ARS investments were intended to provide liquidity through an auction process that resets the applicable interest rate at predetermined calendar intervals, allowing investors to either roll over their holdings or gain immediate liquidity by selling such interests at par. Since February 2008, uncertainties in the credit markets affected our ARS investments and auctions for some of our ARS have continued to fail to settle on their respective settlement dates while some have been redeemed in full at their respective par values. The current portfolio of investments shown as "Long term investments" in our Consolidated Financial Statements represents those investments that are not currently liquid and we will not be able to access these funds until a future auction of these investments is successful, a buyer is found outside of the auction process or the issuer refinances their debt. Maturity dates for these ARS investments range from to 2028 to 2043.

At December 31, 2010, total financial assets measured and recognized at fair value were \$94.8 million and of these assets, \$6.8 million, or 7%, were ARS that were measured and recognized using significant unobservable inputs (Level 3). During 2010, \$650,000 of ARS were redeemed at their full par value, as a result we transferred from Level 3 assets \$465,000 to cash and this resulted in a gain of \$85,000 being recorded to accumulated comprehensive loss in 2010.

As of December 31, 2010, we had \$8.3 million par value (\$6.8 million fair value) of long-term ARS investments. The aggregate loss in value is included as an unrealized loss in accumulated other comprehensive income (loss). Given observable market information was not available to determine the fair values of our ARS portfolio, we valued these investments based on a discounted cash flow model. While our ARS valuation model was based on both Level 2 (credit quality and interest rates) and Level 3 inputs (pricing models), we determined that the Level 3 inputs were the most significant to the overall fair value measurement, particularly the estimates of risk adjusted discount rates. The expected future cash flows of the ARS were discounted using a risk adjusted discount rate that compensated for the illiquidity. Projected future cash flows over the economic life of the ARS (of approximately 7.5 – 15.0 years) were modeled based on the contractual penalty rates for the security added to a tax adjusted LIBOR interest rate curve. The discount rates that were applied to the cash flows were based on a premium over the projected yield curve and included an adjustment for credit, illiquidity, and other risk factors. See Note 1 "Summary of Significant Accounting Policies-Fair Value Measurements" in Notes to Consolidated Financial Statement in Part II, Item 8 of this Form 10-K for more information.

The valuation of our investment portfolio is subject to uncertainties that are difficult to predict. Factors that may impact the valuation include duration of time that the ARS remain illiquid, changes to credit ratings of the securities, rates of default of the underlying assets, changes in the underlying collateral value, market discount rates for similar illiquid investments, and ongoing strength and quality of credit markets. If the auctions for our ARS investments continue to fail, and there is a further decline in their valuation, then we would have to: (i) record additional reductions to the fair value of our ARS investments; and (ii) record unrealized losses in our accumulated comprehensive income (loss) for the losses in value that are associated with market risk. If the decline in fair value is considered other-than-temporary then we would have to record an impairment charge in our Consolidated Statement of Operations for the loss in value associated with the worsening of the credit worthiness (credit losses) of the issuer, which would reduce future earnings and harm our business.

Recognition and Presentation of Other-Than-Temporary-Impairments

We review for impairments on a quarterly basis in order to determine the classification of such as "temporary" or "other-than-temporary." Factors that we consider to make such determination include the duration and severity of the impairment; the reason for the decline in value and the potential recovery period; and our intent to sell, or whether it is more likely than not that we will be required to sell, the investment before recovery. Beginning April 1, 2009, if an entity intends to sell, or if it is more likely than not that we will be required to sell, an impaired debt security prior to recovery of its cost basis, the security is other-than-temporarily impaired and the full amount of the impairment is required to be recognized as a loss through earnings. Otherwise, losses on securities which are other-than-temporarily impaired are separated into:

- (i) the portion of loss which represents the credit loss; or
- (ii) the portion which is due to other factors.

The credit loss portion is recognized as a loss through earnings while the loss due to other factors is recognized in other comprehensive income (loss), net of taxes and related amortization. At December 31, 2010, we had approximately \$8.3 million of par value ARS investments on which we had recognized approximately \$1.5 million in unrealized other-than-temporary losses. Given we believed that such losses were not credit related, we have included them in accumulated comprehensive loss.

Prior to April 1, 2009, all declines in fair value deemed to be other-than-temporary were reflected in earnings as realized losses. With respect to the ARS that we held as of April 1, 2009, we determined that the cumulative effect adjustment required to reclassify the non-credit portion of previously recognized other-than-temporarily impaired adjustments was \$3.5 million. Therefore, we increased our accumulated earnings and decreased our accumulated other comprehensive income (loss) by the \$3.5 million cumulative effect adjustment.

Stock-based Compensation Expense

Employee stock-based compensation is estimated at the date of grant based on the employee stock award's fair value using the Black-Scholes option-pricing model and is recognized as expense ratably over the requisite service period in a manner similar to other forms of compensation paid to employees. The Black-Scholes option-pricing model requires the use of certain subjective assumptions. The most significant of these assumptions are our estimates of the expected volatility of the market price of our stock and the expected term of the award. The expected volatility is a 50%/50% blend of implied and historical volatility. We have determined that this is a more reflective measure of market conditions and a better indicator of expected volatility, than its limited historical volatility since the initial public offering of our common stock. When establishing an estimate of the expected term of an award, we consider historical experience of similar awards, giving consideration to the contractual terms of the awards, vesting requirements, and expectation of future employee behavior, including post-vesting terminations. As required under GAAP, we review our valuation assumptions at each grant date, and, as a result, our valuation assumptions used to value employee stock-based awards granted in future periods may change.

As of December 31, 2010, the unrecognized compensation cost, net of expected forfeitures, was \$5.5 million for stock options and stock awards and \$29,000 for the employee stock purchase plan which will be recognized using the straight-line attribution method over an estimated weighted-average remaining amortization period of 2.40 years and 0.33 years, respectively. See Note 5 "Stockholders' Equity, Stock Plans and Stock-Based Compensation Expense," in the Notes to Consolidated Financial Statement in Part II, Item 8 of this Form 10-K for more information.

Valuation of Inventories

We state our inventories at the lower of cost or market, computed on a standard cost basis, which approximates actual cost on a first-in, first-out basis and market being determined as the lower of replacement cost or net realizable value. Standard costs are monitored and updated quarterly or as necessary, to reflect changes in raw material costs, labor to manufacture the product and overhead rates. We provide for excess and obsolete inventories when conditions indicate that the selling price could be less than cost due to physical deterioration, usage, obsolescence, reductions in estimated future demand and reductions in selling prices. Inventory provisions are measured as the difference between the cost of inventory and estimated market value and charged to cost of revenue to establish a lower cost basis for the inventories. We balance the need to maintain strategic inventory levels with the risk of obsolescence due to changing technology and customer demand levels. Unfavorable changes in market conditions may result in a need for additional inventory provisions that could adversely impact our gross margins. Conversely, favorable changes in demand could result in higher gross margins when product that has previously been reserved is sold.

Warranty Obligations

We historically provided a standard one-year or two-year warranty coverage on our systems. Beginning in September 2009, we changed our warranty policy to a one-year standard warranty on all systems. Warranty coverage provided is for labor and parts necessary to repair the systems during the warranty period. We provide for the estimated future costs of warranty obligations in cost of revenue when the related revenue is recognized. The accrued warranty costs represent our best estimate at the time of sale, and as reviewed and updated quarterly, of the total costs that we expect to incur in repairing or replacing product parts that fail while still under warranty. Accrued warranty costs include costs of material, technical support labor and associated overhead. The amount of accrued estimated warranty costs obligation for established products is primarily based on historical experience as to product failures adjusted for current information on repair costs. Actual warranty costs could differ from the estimated amounts. On a quarterly basis, we review the accrued balances of our warranty obligations and update based on historical warranty cost trends. If we were required to accrue additional warranty cost in the future due to actual product failure rates, material usage, service delivery costs or overhead costs differing from our estimates, revisions to the estimated warranty liability would be required, which would negatively impact our operating results.

Provision for Income Taxes

We are subject to taxes on earnings in both the United States and numerous foreign jurisdictions. As a global taxpayer, significant judgments and estimates are required in evaluating our uncertain tax positions and determining our provision for income taxes on earnings. We perform a two-step approach to recognizing and measuring uncertain tax positions. The first step is to evaluate the tax position for recognition by determining if the weight of available evidence indicates that it is more likely than not that the position will be sustained on audit, including resolution of related appeals or litigation processes, if any. The second step is to measure the tax benefit as the largest amount that is more than 50% likely of being realized upon settlement. Although we believe we have adequately reserved for our uncertain tax positions, no assurance can be given that the final tax outcome of these matters will not be different. We adjust these reserves in light of changing facts and circumstances, such as the closing of a tax audit or the refinement of an estimate. To the extent that the final tax outcome of these matters is different than the amounts recorded, such differences will impact the provision for income taxes in the period in which such determination is made. The provision for income taxes includes the impact of reserve provisions and changes to reserves that are considered appropriate, as well as the related net interest.

Our effective tax rates have differed from the statutory rate primarily due to the tax impact of tax-exempt interest income, foreign operations, research and development tax credits, state taxes, certain benefits realized related to stock option activity, and changes in valuation allowance. Our current effective tax rate does not assume U.S. taxes on undistributed profits of foreign subsidiaries. These earnings could become subject to incremental foreign withholding or U.S. federal and state taxes, should they either be deemed or actually remitted to the United States. The effective tax rate for financial statement provision / (benefit) purposes was approximately 0% in 2010, (102)% in 2009, and 22% in 2008. Our future effective tax rates could be affected by earnings being lower than anticipated in countries where we have lower statutory rates and being higher than anticipated in countries where we have higher statutory rates, or by changes in tax laws, accounting principles, interpretations thereof, net operating loss carryback, research and development tax credits, and due to changes in the valuation allowance of our U.S. deferred tax assets. In addition, we are subject to the examination of our income tax returns by the Internal Revenue Service and other tax authorities. We regularly assess the likelihood of adverse outcomes resulting from these examinations to determine the adequacy of our provision for income taxes.

At December 31, 2010, we had an aggregate of approximately \$3 million of unremitted earnings of foreign subsidiaries that have been, or are intended to be, indefinitely reinvested for continued use in foreign operations. If the total undistributed earnings of foreign subsidiaries were remitted, a significant amount of the additional tax would be offset by the allowable foreign tax credits. It is not practical for us to determine the additional tax of remitting these earnings.

Our deferred tax assets are recognized for the expected future tax consequences of temporary differences between the financial reporting and tax bases of assets and liabilities, and for operating losses and tax credit carryforwards. A valuation allowance reduces deferred tax assets to estimated realizable value, which assumes that it is more likely than not that we will be able to generate sufficient future taxable income in certain tax jurisdictions to realize the net carrying value. The four sources of taxable income to be considered in determining whether a valuation allowance is required include:

- Future reversals of existing taxable temporary differences (i.e., offset gross deferred tax assets against gross deferred tax liabilities);
- Future taxable income exclusive of reversing temporary differences and carryforwards;
- Taxable income in prior carryback years; and
- Tax planning strategies.

Determining whether a valuation allowance for deferred tax assets is necessary requires an analysis of both positive and negative evidence regarding realization of the deferred tax assets. In general, positive evidence may include:

- A strong earnings history exclusive of the loss that created the deductible temporary differences, coupled with evidence indicating that the loss is the result of an aberration rather than a continuing condition; and
- An excess of appreciated asset value over the tax basis of our net assets in an amount sufficient to realize the deferred tax asset.

In general, negative evidence may include:

- A history of operating loss or tax credit carryforwards expiring unused;
- An expectation of being in a cumulative loss position in a future reporting period;
- The existence of cumulative losses in recent years; and
- A carryback or carryforward period that is so brief that it would limit the realization of tax benefits.

The weight given to the potential effect of negative and positive evidence should be commensurate with the extent to which it can be objectively verified and judgment must be used in considering the relative impact of positive and negative evidence.

In evaluating the ability to recover deferred tax assets, we considered available positive and negative evidence, giving greater weight to our recent cumulative losses and our ability to carry-back losses against prior taxable income and lesser weight to its projected financial results due to the challenges of forecasting future periods. We also considered, commensurate with its objective verifiability, the forecast of future taxable income including the reversal of temporary differences. At the end of the quarter ended September 30, 2009, changes in previously anticipated expectations and continued operating losses resulted in a valuation allowance against our tax benefits since we no longer considered them "more-likely-than-not" realizable. We also performed this evaluation as of the year ended December 31, 2010 and determined the full valuation allowance was still required.

Long-Lived Asset Impairment

Long-lived assets, such as property and equipment and intangible assets, are reviewed for impairment whenever events or changes in circumstances indicate that the carrying amount of such assets may not ultimately be recoverable. Determination of recoverability is based on an estimate of undiscounted future cash flows resulting from the use of the asset and its ultimate disposition. If the sum of the expected future cash flows is less than the carrying amount of those assets, we recognize an impairment loss based on the excess of the carrying amount over the fair value of the assets. Through December 31, 2010, there have been no such impairments.

Litigation

We have been, and may in the future become, subject to legal proceedings related to securities litigation, intellectual property and other matters such as the TCPA litigation and the securities class Action Lawsuit described in Item 3—Legal Proceedings. Based on all available information at the balance sheet dates, we assess the likelihood of any adverse judgments or outcomes for these matters, as well as potential ranges of probable loss. If losses are probable and reasonably estimable, we record a reserve. See "Item 3 - Legal Proceedings" in Part I, of this Form 10-K.

Recent Accounting Guidance

For a full description of recent accounting pronouncements, including the respective expected dates of adoption and effects on results of operations and financial condition see Note 1 "Summary of Significant Accounting Policies — New Accounting Standards" in the Notes to Consolidated Financial Statements in Part II, Item 8 of this Form 10-K.

Results of Operations

The following table sets forth selected consolidated financial data expressed as a percentage of net revenue.

	Year Ended December 31,					
	2010	2009	2008			
Net revenue	100%	100%	100%			
Cost of revenue	43%	41%	39%			
Gross profit	57%	59%	61%			
Operating expenses:						
Sales and marketing	47%	45%	42%			
Research and development	13%	13%	9%			
General and administrative	18%	19%	14%			
Litigation settlement	%	1%	%			
Total operating expenses	78%	78%	65%			
Loss from operations	(21)%	(19)%	(4)%			
Interest and other income, net	1%	3%	4%			
Other-than-temporary impairment of long-term investments	<u> </u>	%	(4)%			
Loss before income taxes	(20)%	(16)%	(4)%			
Provision (benefit) for income taxes	%	17%	(1)%			
Net loss	(20)%	(33)%	(3)%			

Net Revenue

The following table sets forth selected consolidated revenue by major geographic area and product category with changes thereof.

	Year Ended December 31,										
(Dollars in thousands)	2010	% Change	2009(1)	% Change	2008(1)						
Revenue mix by geography:											
United States	\$ 19,337	(8)% \$	21,019	(50)%\$	41,683						
Percent of total	36%	· · · ·	39%	\	50%						
Japan	\$ 13,625	41% \$	9,636	(12)%\$	10,929						
Asia, excluding Japan	5,131	9%	4,727	(17)%	5,713						
Europe	5,801	(18)%	7,087	(33)%	10,522						
Rest of the world	9,380	(16)%	11,213	(23)%	14,532						
Total international revenue	33,937	4%	32,663	(22)%	41,696						
Percent of total	64%	_	61%	` ′ –	50%						
Total consolidated revenue	\$ 53,274	(1)% <u>\$</u>	53,682	(36)% <u>\$</u>	83,379						
Revenue mix by product category:											
Products	\$ 27,808	4% \$	26,842	(54)%\$	57,821						
Upgrades	4,824	(24)%	6,343	(24)%	8,361						
Service	13,231	<u>_</u> %	13,186	16%	11,358						
Titan hand piece refills	3,863	(31)%	5,599	(1)%	5,662						
Dermal fillers and cosmeceuticals ⁽¹⁾	3,548	ì07%	1,712	867%	177						
Total consolidated revenue	\$ 53,274	(1)% <u>\$</u>	53,682	(36)% \(\bar{\bar{\bar{\bar{\bar{\bar{\bar{	83,379						

⁽¹⁾ Beginning in 2010, we classified revenue from dermal fillers and cosmeceuticals product in the revenue category 'Dermal fillers and cosmeceuticals.' Previously, we classified this revenue under the category of 'Products.' As such, we reclassified the 2009 and 2008 revenue from 'Products' to 'Dermal fillers and cosmeceuticals.'

Revenue by Geography:

In 2010 our net revenue declined by 1%, compared to 2009, and in 2009 it declined by 36%, compared to 2008.

Our U.S. revenue decreased by 8% in 2010, compared to 2009, and by 50% in 2009, compared to 2008. We believe the decline in U.S. revenues was attributable to several factors, including:

- Our Products and Upgrades ASPs declined in 2010 and 2009, compared to their respective prior years.
 This was attributable primarily to customers purchasing fewer applications for systems and lower pricing resulting from competitive discounting pressures.
- Though our unit sales of Products and Upgrades increased in 2010, compared with 2009, they declined in 2009, compared to 2008.
- We experienced a temporary decline in our Titan refill revenue in 2010, compared to 2009, due to a voluntary recall of certain Titan XL hand pieces. All customers that had a Titan XL hand pieces subject to the recall, were provided with fully "refilled" hand pieces, which delayed their purchase of a refill.

International revenues increased by 4% in 2010, compared to 2009, and decreased by 22% in 2009, compared to 2008. The growth in our international revenue in 2010 was sourced from most of the international locations that we sell in. In Japan, our revenue increased by 41% in 2010, compared to 2009, as a result of growth from all of our product categories and the recently added Dermal fillers and cosmeceuticals business. With respect to Australia, our revenue declined in 2010 by approximately 65% due to 2009 being an unusually high revenue year in which a special tax incentive was offered by the government to incentivize customers to purchase capital equipment. The 2009 decline in our international revenue, compared to 2008, was primarily attributable to the global recession that caused our current and prospective customers to be reluctant to spend significant amounts of money on capital equipment purchases during the unstable economic time.

Revenue by Product Category:

Our product revenue increased by 4% in 2010, compared to 2009, and decreased by 54% in 2009, compared to 2008. The 2010 increase in product revenue was primarily attributable to revenue from the GenesisPlus product that was launched in the third quarter of 2010. The 2009 declines in Product revenue, compared to 2008, was primarily attributable to the global recession that caused our current and prospective customers to be reluctant to spend significant amounts of money on capital equipment purchases during the unstable economic times. We believe that in 2009 and in 2010, some of our U.S. current and prospective customers that did not have established medical offices, continued to be reluctant to purchase capital equipment due to the general economic uncertainty and tight credit conditions.

Upgrade revenue decreased by 24% in both 2010 and 2009, compared to the respective prior year periods. In the past, we introduced new products that allowed existing customers to upgrade their previously purchased systems to take benefit of the additional capabilities. However, in 2010 we introduced GenesisPlus- a stand alone product- and in 2009 we did not have a new product introduction, which resulted in the decline in our Upgrade revenue. Further, our ASPs for Upgrades declined in 2010 and 2009, compared to the respective prior year periods, as a result of customers purchasing fewer handpieces and due to competitive discounting pressures.

Our Service revenue remained relatively flat in 2010, compared to 2009, and increased by 16% in 2009, compared to 2008. Service contract amortization is the primary component of our service revenue. In years prior to 2009, we had discounted pricing promotions for incentivizing customers to purchase multiple year service contracts, however, in 2009 and 2010 we did not have such promotions.

As a result, our service revenue increased in 2009, compared to 2008, due to the continuing amortization of deferred service contracts purchased prior to 2009. However, in 2010, service revenue remained flat as a result of the decline in unit sales in 2009 that included an element of deferred revenue for service contracts beyond our standard one-year warranty term.

Our Titan hand piece refill revenue decreased 31% in 2010, compared to 2009, and decreased 1% in 2009, compared to 2008. The decrease in 2010 was due to a voluntary recall of certain Titan XL hand pieces whereby eligible customers were provided with fully refilled hand pieces.

Our Dermal filler and cosmeceutical business increased by 107% in 2010, compared to 2009. This increase was attributable primarily due to the commencement in 2010 of the distribution of Obagi's cosmeceutical products to physicians in the Japanese market.

Gross Profit

	Year Ended December 31,										
(Dollars in thousands)	 2010	% Change	2009	% Change		2008	٠.				
Gross Profit	\$ 30,216	(5)% \$	31,923	(37)%	\$	51,021	٠				
As a percentage of total revenue	57%		599	6:		619	6				

Our cost of revenue consists primarily of materials, personnel expenses, royalty expense, warranty and manufacturing overhead expenses. Gross margin as a percentage of net revenue was 57% in 2010, 59% in 2009 and 61% in 2008. We believe the decrease in gross margins in 2010, compared to 2009, and 2009, compared to 2008, was primarily attributable to the following:

- The 2010 voluntary recall of certain Titan XL hand pieces whereby eligible customers were provided with fully refilled hand pieces;
- A \$1.7 million, or 31%, temporary decrease in our Titan refill revenue in 2010, compared to 2009, for which we traditionally earn a higher gross margin than our blended total gross margin percentage;
- Our ASPs have declined due primarily to customers purchasing fewer applications on their platforms and due to competitive discounting pressures;
- A higher proportion of distributor revenue that carries a lower gross margin; partially offset by
- Lower manufacturing expenses resulting from headcount reductions; and
- Reduced warranty and service expenses as a result of improved product reliability (for products other than Titan XL hand pieces).

Sales and Marketing

	Year Ended December 31,										
(Dollars in thousands)	٠.	2010	% Change		2009	% C	hange		2008		
Sales and marketing	\$	24,735	2%	\$	24,286		(31)%	\$	35,354		
As a percentage of total revenue		47%	to a service of		45%	:			42%		
		100	The second of the second	45	· · · ·						

Our sales and marketing expenses consist primarily of personnel expenses, expenses associated with customer-attended workshops and trade shows, and advertising. Sales and marketing expenses increased \$449,000 in 2010, compared to 2009, which was primarily attributable to the following:

- \$855,000 increase in personnel expenses in marketing due primarily to an increase in headcount resulting from the creation of three new departments: post marketing studies (clinical development), business development and telesales;
- \$242,000 increase in international spending on workshops, advertising and other promotional activities; offset by
- A decline in U.S. sales personnel expenses by \$617,000 due to lower headcount; and due to decreased sales commissions resulting from lower U.S revenue.

In 2009 sales and marketing expenses decreased by \$11.1 million, compared to 2008. This decrease was primarily attributable to:

- lower salaries and commissions expenses of \$5.4 million as a result of our reduction in force during the first half of 2009 and lower sales volumes in 2009;
- lower travel and travel related expenses of \$1.8 million; and
- lower marketing expenses of \$983,000 as a result of less spending on workshops, advertising and other promotional activities.

Sales and marketing expenses as a percentage of total revenue, increased to 46% in 2010, compared to 45% in 2009 and 42% in 2008. This increase was due primarily to a decline in our total revenue in 2009 and 2010.

Research and Development (R&D)

(Dollars in thousands)		2010	% Change	2009	% Change		2008
Research and development	\$	7,004	3%	\$ 6,810	(10)%	\$	7,550
As a percentage of total revenue		13%		13%		1	9%

Research and development (R&D) expenses consist primarily of personnel expenses, clinical, regulatory and material costs. R&D expenses increased \$194,000 in 2010, compared to 2009, which was due primarily to higher personnel expenses resulting from higher headcount in engineering relating to new product development programs.

In 2009 R&D expenses decreased by \$740,000, compared to 2008, which was due primarily to lower headcount (partially resulting from a reduction-in-force that we implemented in the first-half of 2009) and a reduction in consulting services of \$689,000. R&D expenses as a percentage of total revenue, increased to 13% in 2009, compared to 9% in 2008, due primarily to a function of lower revenue in 2009 mitigated by cost containment.

General and Administrative (G&A)

· · ·	Year Ended December 31,									
(Dollars in thousands)	* -	2010	% Change		2009	% Change		2008		
General and administrative	\$	9,576	(7)%	\$	10,320	(8)%	\$	11,270		
As a percentage of total revenue		18%			19%			14%		

General and administrative expenses consist primarily of: personnel expenses, legal fees, accounting, audit and tax consulting fees, and other general and administrative expenses. G&A expenses decreased by \$744,000 in 2010, compared to 2009, which was primarily attributable to:

- a \$626,000 reduction in bad debts expense due to a large non recurring expense in 2009; and
- a \$201,000 reduction in legal fees and legal settlement expenses.

In 2009 G&A expenses decreased by \$950,000, compared to 2008. This decrease was primarily attributable to:

- a decrease in legal, audit and tax consulting fees of \$587,000, due to reduced fees from the consulting firms, partially offset by higher consulting fees related to our 2009 Option Exchange Program;
- a decrease in personnel expenses of \$206,000 due primarily to lower headcount (resulting from a reduction-in-force that we implemented in the first-half of 2009); partly offset by
- an increase in bad debt expense of \$392,000, resulting primarily from one leasing company that defaulted on its payment in the second quarter of 2009 due to it having significant financial problems.

Litigation Settlement

In 2009, we settled our TCPA class action lawsuit and in that regard recorded a charge of \$850,000 for the cost of the settlement, net of administrative expenses and amounts that were recovered from our insurance carrier.

Interest and Other Income, Net

The components of "Interest and Other Income, Net" are as follows:

	Year Ended December 31,											
(Dollars in thousands)		2010	% Change		2009	% C	hange		2008			
Interest income	\$	539	(61)%	\$	1,383	· <u>·</u>	(56)%	\$	3,170			
Other income (expense), net		44	(77)%		189	1	NA		(124)			
Total Interest and other income, net	\$	583	(63)%	\$	1,572		(48)%	\$	3,046			

Interest income decreased 61% in 2010, compared to 2009, and decreased 56% in 2009, compared to 2008. These decreases were due primarily to reduced tax-exempt interest yields, as a result of lower interest rates, and reduced investment balance. Our cash, cash equivalents, marketable investments and long-term investments measured and recognized at fair value were \$96.8 million at December 31, 2010, \$106.9 million at December 31, 2009 and \$106.8 million December 31, 2008.

Other-Than-Temporary Impairments of Long-Term Investments

		·			
(Dollars in thousands)	2010	% Change	2009	% Change	2008
Other-than-temporary impairment of long-			$(x,y) = (x,y) \cdot (x,y)$		
term investments	\$ —	NA	\$ —	(100)%	\$ 3,554

For the year ended December 31, 2008, we determined there was a decline in the fair value of our ARS investments for which we recorded a \$3.6 million other-than-temporary impairment charge. See the 'Critical Accounting Estimates' section above, for additional details relating to the charge.

Provision (Benefit) for Income Taxes

	Year Ended December 31,											
(Dollars in thousands)	2010	\$	Change		2009	\$	Change		2008			
Loss before income taxes	\$ (10,516)	\$	(1,745)	\$	(8,771)	\$	(5110)	\$	(3,661)			
Provision (benefit) for income taxes	2		(8,906)	,	8,908		9,700		(792)			
Effective tax rate	09	6	-		(102)%	ó			22%			

We recognized a \$2,000 income tax provision in 2010, resulting in a 0% effective tax rate, and due principally to a full valuation allowance applied against deferred tax assets arising during the year. We recognized an income tax provision of \$8.9 million in 2009, despite losses before taxes. The provision was primarily due to the recording of a valuation allowance at the end of the third quarter of 2009 to reduce certain U.S. federal and state net deferred tax assets to their anticipated realizable value, of which \$10.2 million related to our U.S. deferred tax assets as of December 31, 2008. This valuation allowance was offset by \$1.3 million of certain tax benefits resulting from losses generated during fiscal 2009 that were carried-back to prior periods.

ASC 740 requires the consideration of a valuation allowance to reflect the likelihood of realization of deferred tax assets. Significant management judgment is required in determining any valuation allowance recorded against deferred tax assets. In evaluating the ability to recover deferred tax assets we considered available positive and negative evidence, giving greater weight to our recent cumulative losses and our ability to carry-back losses against prior taxable income and lesser weight to our projected financial results due to challenges of forecasting future periods. We also considered, commensurate with its objective verifiability, the forecast of future taxable income including the reversal of temporary differences. At the end of the quarter ended September 30, 2009, revisions in previously anticipated expectations and continued operating losses resulted in a valuation allowance against our tax benefits since it was not considered "more-likely-than-not" realizable. We also performed this evaluation as of the years ended December 31, 2009 and 2010 and determined the full valuation allowance was still required. Under current tax laws, this valuation allowance will not limit our ability to utilize federal and state deferred tax assets provided we can generate sufficient future taxable income in the U.S.

We anticipate we will continue to record a valuation allowance against the losses of certain jurisdictions, primarily U.S. federal and state, until such time as we are able to determine it is "more-likely-than-not" the deferred tax asset will be realized. Such position is dependent on whether there will be sufficient future taxable income to realize such deferred tax assets. We expect our future tax provisions (benefits), during the time such valuation allowances are recorded, will consist primarily of the tax expense of our non-U.S. jurisdictions that are profitable. Our effective tax rate may vary from period to period based on changes in estimated taxable income or loss by jurisdiction, changes to the valuation allowance, changes to federal, state or foreign tax laws, future expansion into areas with varying country, state, and local income tax rates, deductibility of certain costs and expenses by jurisdiction.

Net Loss and Net Loss per Diluted Share

	Year Ended December 31,									
(Dollars in thousands, except per share data)		2010	% Change	2009	% Change		2008			
Net loss	\$	(10,518)	(41)% \$	(17,679)	516%	<u>`</u>	(2,869)			
Net loss per diluted share	\$	(0.78)	(41)% \$	(1.33)	505% \$	3	(0.22)			

The \$7.2 million decrease in net loss, and \$0.55 decrease in net loss per diluted share in 2010, compared to 2009, was primarily attributable to:

- a reduction in the tax provision by \$8.9 million, which was principally a function of the deferred tax asset valuation allowances recorded in 2009;
- lower operating expenses of \$951,000, due primarily to the \$850,000 litigation settlement expense in 2009 not being incurred in 2010;
- offset by a decline in our gross profit by \$1.7 million and other income by \$989,000.

The \$14.8 million increase in net loss, and \$1.11 increase in net loss per diluted share in 2009, compared to 2008, was primarily attributable to lower revenue of \$29.7 million, partially offset by a decrease of \$11.9 million in operating expenses.

Liquidity and Capital Resources

Liquidity is the measurement of our ability to meet potential cash requirements, fund the planned expansion of our operations and acquire businesses. Our sources of cash include operations, stock option exercises, and employee stock purchases. We actively manage our cash usage and investment of liquid cash to ensure the maintenance of sufficient funds to meet our daily needs. The majority of our cash and investments are held in U.S. banks and our foreign subsidiaries maintain a limited amount of cash in their local banks to cover their short-term operating expenses. The following table summarizes our cash and cash equivalents, marketable investments and long-term investments (in thousands):

			As of December 31,						
(Dollars in thousands)		2010		2009		Change			
Cash, cash equivalents and marketable securities:									
Cash and cash equivalents	\$	12,519	\$	22,829	\$	(10,310)			
Marketable investments		77,484		76,780		704			
Long-term investments		6,784		7,275		(491)			
Total	\$	96,787	\$	106,884	\$	(10,097)			

Cash Flows

In summary, our cash flows were as flows:

	Year ended December 31,							
(Dollars in thousands)		2010 20				2008		
Cash flows provided by (used in):								
Operating activities	\$	(8,059)	\$	41	\$	4,340		
Investing activities		(2,777)		(14,360)		20,644		
Financing activities		526		608		502		
Net increase (decrease) in cash and cash equivalents	\$	(10,310)	\$	(13,711)	\$	25,486		

Cash Flows from Operating Activities

We used net cash of \$8.1 million in operating activities during 2010, which was primarily attributable to:

- \$5.2 million used from net loss of \$10.5 million after adjusting for non-cash related items of \$5.3 million, consisting primarily of stock based compensation expense of \$4.7 million and depreciation and amortization expense of \$717,000;
- \$2.6 million used as a result of a decrease in accrued liabilities due primarily to a reduction in the liability for warranty costs of \$253,000 resulting primarily from a reduction in the total units remaining under warranty, a decrease in accrued expenses of \$1.4 million for payroll, professional services, sales & marketing, and other miscellaneous expenses resulting from continued cost containment initiatives, and a reduction of approximately \$950,000 for the pay out of our prior year accrual for the TCPA class action lawsuit; and
- \$1.2 million used as a result of a decrease in deferred revenue due primarily to a reduction in deferred service contracts resulting from a decline in our sales unit volume in 2009 and a reduction in the pricing charged for service contracts; partially offset by
- \$2.3 million generated from a reduction in other current assets and prepaid expenses, resulting primarily from a reduction in accrued interest and unamortized discounts related to our marketable and long-term investments.

We generated net cash from operating activities of \$41,000 in 2009, which was primarily attributable to:

- \$849,000 used from net loss of \$17.7 million after adjusting for non-cash related items of \$16.8 million, consisting primarily of a valuation allowance on our deferred tax asset of \$10.5 million, stock-based compensation expense of \$4.2 million, net increase in the allowance for doubtful accounts of \$525,000 due primarily to one leasing company that has defaulted on its payment, and an increase in the provision for excess and obsolete inventories of \$611,000 resulting from the reduced future demand for our products; and
- \$3.5 million used as a result of a decrease in deferred revenue due primarily to a decrease in unit sales
 volume of Products and Upgrades that included purchases of extended service contracts, a reduction in
 our service contract pricing beginning in 2010, a shift by customers towards purchasing shorter term
 contracts, and fewer customers purchasing extended service contracts in response to improved product
 reliability and to a tougher economy; offset by
- \$2.9 million of cash generated by the decrease in gross inventory balance from December 31, 2009 to December 31, 2010, that resulted from slowing our inventory build to better match the reduced sales of our products; and
- \$1.9 million of cash generated by the decrease in gross accounts receivable balance from December 31, 2009 to December 31, 2010 that resulted from the collection of the higher 2009 year-end accounts receivable balances.

Cash Flows from Investing Activities

We used net cash of \$2.8 million from investing activities in 2010, which was primarily attributable to:

- \$85.3 million in net proceeds from the sales and maturities of \$650,000 of our ARS investments and due to us diversifying out of municipal securities into other secure financial instruments; partially offset by
- \$87.8 million of cash used to purchase marketable investments; and
- \$275,000 of cash used to purchase property and equipment.

We used net cash of \$14.4 million from investing activities in 2009, which was primarily attributable to:

- \$53.7 million of cash used to purchase marketable investments; partially offset by
- \$39.4 million in net proceeds from the sales and maturities of marketable investments.

Cash Flows from Financing Activities

Net cash provided by financing activities in 2010 was \$526,000, which resulted from \$518,000 of cash generated by the issuance of stock through our stock option and employee stock purchase plans and \$8,000 of excess tax benefits related to stock-based compensation expenses reclassified from operating activities to financing activities.

Net cash provided by financing activities in 2009 was \$608,000, which resulted from \$585,000 of cash generated by the issuance of stock through our stock option and employee stock purchase plans and \$23,000 of excess tax benefits related to stock-based compensation expenses reclassified from operating activities to financing activities in accordance with FAS 123(R).

Adequacy of cash resources to meet future needs

We had cash, cash equivalents, marketable and long-term investments of \$96.8 million as of December 31, 2010. Of this amount, we had \$6.8 million invested in long-term ARS investments (see 'Critical Accounting Policies and Estimates' section above, for a full description of our long-term investments in ARS). We believe that our existing cash resources are sufficient to meet our anticipated cash needs for working capital and capital expenditures for at least the next 12 months.

Off-Balance Sheet Arrangements

We do not participate in transactions that generate relationships with unconsolidated entities or financial partnerships, such as entities often referred to as structured finance, variable interest or special purpose entities, which would have been established for the purpose of facilitating off-balance sheet arrangements or other contractually narrow or limited purposes.

We have certain contractual arrangements that create potential risk for us and are not recognized in our Consolidated Balance Sheets. Discussed below are off-balance sheet arrangements that have or are reasonably likely to have a material current or future effect on our financial condition, changes in financial condition, revenue or expenses, results of operations, liquidity, capital expenditures, or capital resources.

We lease various real properties under operating leases that generally require us to pay taxes, insurance, maintenance, and minimum lease payments. Some of our leases have options to renew.

Contractual Obligations

The following are our obligations for future minimum lease commitments related to facility leases as of December 31, 2010:

the state of the s									
•	 Less Than						-	Mo	re Than
Contractual Obligations	 Total	1	Year	1-3	3 Years	3-	5 Years	5 Years	
Operating leases	\$ 9,632	\$	1,688	\$ 2,790		\$	2,501	\$	2,653

Purchase Commitments

We maintain certain open inventory purchase commitments with our suppliers to ensure a smooth and continuous supply for key components. Our liability in these purchase commitments is generally restricted to a forecasted time-horizon as agreed between the parties. These forecasted time-horizons can vary among different suppliers. Our open inventory purchase commitments were not material at December 31, 2010. As a result, this amount is not included in the contractual obligations table above.

Income Tax Liability

We have included in our Consolidated Balance Sheet \$477,000 in long-term income tax liability with respect to unrecognized tax benefits and accrued interest as of December 31, 2010. At this time, we are unable to make a reasonably reliable estimate of the timing of payments in individual years beyond 12 months due to uncertainties in the timing of tax audit outcomes. As a result, this amount is not included in the contractual obligations table above.

Other

In the normal course of business, we enter into agreements that contain a variety of representations, warranties, and indemnification obligations. For example, we have entered into indemnification agreements with each of our directors and executive officers. Our exposure under the various indemnification obligations is unknown and not reasonably estimable as they involve future claims that may be made against us. As such, we have not accrued any amounts for such obligations.

ITEM 7A. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK

Interest Rate Sensitivity

Our exposure to interest rate risk relates primarily to our investment portfolio. Fixed rate securities may have their fair market value adversely impacted due to fluctuations in interest rates, while floating rate securities may produce less income than expected if interest rates fall. Due in part to these factors, our future investment income may fall short of expectation due to changes in interest rates or we may suffer losses in principal if forced to sell securities which have declined in market value due to changes in interest rates. The primary objective of our investment activities is to preserve principal while at the same time maximizing yields without significantly increasing risk. To achieve this objective, we invest in debt instruments of the U.S. Government and its agencies and municipal bonds, and, by policy, restrict our exposure to any single type of investment or issuer by imposing concentration limits. To minimize the exposure due to adverse shifts in interest rates, we maintain investments at a weighted average maturity (interest reset date for ARS) of generally less than eighteen months. Assuming a hypothetical increase in interest rates of one percentage point, the fair value of our total investment portfolio would have potentially declined by approximately \$651,000 as of December 31, 2010.

We hold interest bearing ARS that represent investments in pools of student loans issued by the Federal Family Education Loan Program. At the time of acquisition, these ARS investments were intended to provide liquidity via an auction process that resets the applicable interest rate at predetermined calendar intervals, allowing investors to either roll over their holdings or gain immediate liquidity by selling such interests at par. Since February 2008, uncertainties in the credit markets affected our holdings in ARS investments and auctions for all of our investments in these securities failed until December 31, 2008. In 2010 and 2009, approximately \$650,000 and \$4.4 million, respectively of our original \$13.4 million par value portfolio has been redeemed in full and as of December 31, 2010 we had \$8.3 million par value (fair value of \$6.8 million) of long-term ARS, whose auctions continue to fail. These investments are not currently liquid and we will not be able to access these funds until a future auction of these investments is successful, a buyer is found outside of the auction process or the ARS is refinanced by the issuer into another type of debt instrument. Maturity dates for these ARS investments range from 2028 to 2043. We currently classify all of these investments as long-term investments in our Consolidated Balance Sheet because of our continuing inability to determine when these investments will settle. We have also modified our current investment strategy and increased our investments in more liquid money market investments, United States Treasury securities, municipal bonds, and eliminated investments in corporate debt. The valuation of our ARS investment portfolio is subject to uncertainties that are difficult to predict. Factors that may impact its valuation include, duration of time that the ARS remain illiquid, changes to credit ratings of the securities, rates of default of the underlying assets, changes in the underlying collateral value, market discount rates for similar illiquid investments, ongoing strength and quality of credit markets. If the auctions for our ARS investments continue to fail, and there is a further decline in the valuation, then we would have to: (i) record additional reductions to the fair value of our ARS investments; and (ii) record unrealized losses in our accumulated comprehensive income (loss) for the losses in value that are associated with market risk. If the decline in fair value is considered other-than-temporary then we would have to record an impairment charge in our Consolidated Statement of Operations for the loss in value associated with the worsening of the credit worthiness (credit losses) of the issuer, which would reduce future earnings and harm our business.

Foreign Currency Exchange Risk

We have international subsidiaries and operations and are, therefore, subject to foreign currency rate exposure. Although the majority of our revenue and purchases are denominated in U.S. dollars, we have revenue to certain international customers and expenses denominated in the Japanese Yen, Euro, Pounds Sterling, Australian Dollars, Swiss Francs and Canadian Dollars. The net gains and losses from the revaluation of foreign denominated assets and liabilities was a loss of approximately \$34,000 in 2010, which is included in our Consolidated Statements of Operations. Movements in currency exchange rates could cause variability in our revenues, expenses or interest and other income (expense). Though to date our exposure to exchange rate volatility has not been significant, we cannot assure that there will not be a material impact in the future. Future fluctuations in the value of the U.S. dollar may affect the price competitiveness of our products. We do not believe, however, that we currently have significant direct foreign currency exchange rate risk and have not hedged exposures denominated in foreign currencies.

We do not utilize derivative financial instruments, derivative commodity instruments or other market risk sensitive instruments.

ITEM 8. FINANCIAL STATEMENTS AND SUPPLEMENTARY DATA

CUTERA, INC. AND SUBSIDIARY COMPANIES

ANNUAL REPORT ON FORM 10-K

INDEX TO CONSOLIDATED FINANCIAL STATEMENTS

The following Consolidated Financial Statements of the Registrant and its subsidiaries are required to be included in Item 8:

Report of Independent Registered Public Consolidated Balance Sheets					
Consolidated Statements of Operations					
Consolidated Statements of Stockholders			1.	* , *	
Consolidated Statements of Cash Flows.		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	7 () 2 ()	58
Notes to Consolidated Financial Statemer	nts			•	59

The following Consolidated Financial Statement Schedule of the Registrant and its subsidiaries for the years ended December 31, 2010, 2009 and 2008 is filed as a part of this Report as required to be included in Item 15(a) and should be read in conjunction with the Consolidated Financial Statements of the Registrant and its subsidiaries:

Schedule		Page
II	Valuation and Qualifying Accounts	82

All other required schedules are omitted because of the absence of conditions under which they are required or because the required information is given in the Consolidated Financial Statements or the Notes thereto.

Report Of Independent Registered Public Accounting Firm

To the Board of Directors and Stockholders of Cutera, Inc.:

In our opinion, the consolidated financial statements listed in the accompanying index present fairly, in all material respects, the financial position of Cutera, Inc. and its subsidiaries at December 31, 2010 and December 31, 2009, and the results of their operations and their cash flows for each of the three years in the period ended December 31, 2010 in conformity with accounting principles generally accepted in the United States of America. In addition, in our opinion, the financial statement schedule listed in the accompanying index presents fairly, in all material respects, the information set forth therein when read in conjunction with the related consolidated financial statements. Also in our opinion, the Company maintained, in all material respects, effective internal control over financial reporting as of December 31, 2010, based on criteria established in Internal Control - Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission (COSO). The Company's management is responsible for these financial statements and financial statement schedule, for maintaining effective internal control over financial reporting and for its assessment of the effectiveness of internal control over financial reporting, included in Management's Report on Internal Control over Financial Reporting under Item 9A. Our responsibility is to express opinions on these financial statements, on the financial statement schedule, and on the Company's internal control over financial reporting based on our integrated audits. We conducted our audits in accordance with the standards of the Public Company Accounting Oversight Board (United States). Those standards require that we plan and perform the audits to obtain reasonable assurance about whether the financial statements are free of material misstatement and whether effective internal control over financial reporting was maintained in all material respects. Our audits of the financial statements included examining, on a test basis, evidence supporting the amounts and disclosures in the financial statements, assessing the accounting principles used and significant estimates made by management, and evaluating the overall financial statement presentation. Our audit of internal control over financial reporting included obtaining an understanding of internal control over financial reporting, assessing the risk that a material weakness exists, and testing and evaluating the design and operating effectiveness of internal control based on the assessed risk. Our audits also included performing such other procedures as we considered necessary in the circumstances. We believe that our audits provide a reasonable basis for our opinions.

As discussed in Note 1 to the consolidated financial statements, the Company changed the manner in which it accounts for other-than-temporary impairments in 2009.

A company's internal control over financial reporting is a process designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles. A company's internal control over financial reporting includes those policies and procedures that (i) pertain to the maintenance of records that, in reasonable detail, accurately and fairly reflect the transactions and dispositions of the assets of the company; (ii) provide reasonable assurance that transactions are recorded as necessary to permit preparation of financial statements in accordance with generally accepted accounting principles, and that receipts and expenditures of the company are being made only in accordance with authorizations of management and directors of the company; and (iii) provide reasonable assurance regarding prevention or timely detection of unauthorized acquisition, use, or disposition of the company's assets that could have a material effect on the financial statements.

Because of its inherent limitations, internal control over financial reporting may not prevent or detect misstatements. Also, projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because of changes in conditions, or that the degree of compliance with the policies or procedures may deteriorate.

/s/ PricewaterhouseCoopers

San Jose, California March 15, 2011

CUTERA, INC. CONSOLIDATED BALANCE SHEETS (in thousands, except share and per share data)

	 Decem	ber 31,		
	2010		2009	
Assets			Çarin e	
Current assets:				
Cash and cash equivalents	\$ 12,519	\$	22,829	
Marketable investments	77,484		76,780	
Accounts receivable, net of allowance for doubtful accounts in 2010 and				
2009 of \$20 and \$586, respectively	 4,208		3,327	
Inventories	6,448		6,408	
Deferred tax asset	63	,	175	
Other current assets and prepaid expenses	2,740		2,785	
Total current assets	 103,462		112,304	
Property and equipment, net	597		847	
Long-term investments	 6,784		7,275	
Intangibles, net	637		829	
Deferred tax asset, net of current portion	325	4.	97	
Total assets	\$ 111,805	\$	121,352	
Liabilities and Stockholders' Equity				
Current liabilities:				
Accounts payable	\$ 1,296	\$	1.081	
Accrued liabilities	6,194	,	9,048	
Deferred revenue	5,633		6,160	
Total current liabilities	 13,123		16,289	
Deferred rent	1,501		1,493	
Deferred revenue, net of current portion	1,287		1,968	
Income tax liability	477		749	
Total liabilities	16,388		20,499	
Commitments and contingencies (Note 11)	 			
Stockholders' equity:				
Convertible preferred stock, \$0.001 par value Authorized: 5,000,000 shares; none				
issued and outstanding				
Common stock, \$0.001 par value:				
Authorized: 50,000,000 shares; Issued and outstanding: 13,629,713 and				
13,436,163 shares in 2010 and 2009, respectively	14		13	
Additional paid-in capital	90,423		85,248	
Retained earnings	6,736		17,254	
Accumulated other comprehensive loss	(1,756)		(1,662)	
Total stockholders' equity	95,417		100,853	
Total liabilities and stockholders' equity	\$ 111,805	\$	121,352	

CUTERA, INC. CONSOLIDATED STATEMENTS OF OPERATIONS (in thousands, except per share data)

$a^{(1)} = a^{(2)} + a^{(2)}$	Year Ended December 31,						
		2010		2009		2008	
Net revenue	\$	53,274	\$	53,682	\$	83,379	
Cost of revenue		23,058		21,759		32,358	
Gross profit		30,216		31,923		51,021	
Operating expenses:							
Sales and marketing		24,735		24,286		35,354	
Research and development		7,004		6,810		7,550	
General and administrative		9,576		10,320		11,270	
Litigation settlement		_		850			
Total operating expenses	_	41,315		42,266		54,174	
Loss from operations		(11,099)		(10,343)		(3,153)	
Interest and other income, net		583		1,572		3,046	
Other-than-temporary impairments of long-term investments		<u> </u>				(3,554)	
Loss before income taxes		(10,516)		(8,771)		(3,661)	
Provision (benefit) for income taxes		2		8,908		(792)	
Net loss	\$	(10,518)	\$	(17,679)	\$	(2,869)	
Net loss per share:		•					
Basic and diluted	\$	(0.78)	\$	(1.33)	\$	(0.22)	
Weighted-average number of shares used in per share calculations:							
Basic and diluted		13,540		13,279		12,770	

The accompanying notes are an integral part of these consolidated financial statements.

CUTERA, INC. CONSOLIDATED STATEMENTS OF STOCKHOLDERS' EQUITY (in thousands, except share amounts)

	Commor	ı Stock	Additional Paid-in	Deferred Stock-Based	Retained	Accumulated Other Comprehensive	Total Stockholders'
	Shares	Amount	Capital	Compensation	Earnings	Income (loss)	Equity
Balance at December 31,							
2007 Issuance of common stock for employee purchase	12,738,449	13	74,871	· . —	34,279	190	109,353
plan	50,693		464	_	_	_	464
Exercise of stock options Issuance of common stock in settlement of restricted stock units, net of shares withheld for employee	8,449	_	45	_	_	_	45
taxes	8,444	_	(51)	_	_	-	(51)
expense Tax benefit from exercises of stock-based payment	_	_	5,220	_	_	_	5,220
awards		_	(231)	_	_	_	(231)
Net loss Other comprehensive	_	_	_	_	(2,869)	_	(2,869)
income, net of tax of \$230 Comprehensive	_	_	·—.	<u> </u>	_	177	177
loss	_		<u>. </u>			_	(2,692)
Balance at December 31, 2008	12,806,035	\$ 13	\$ 80,318	<u> </u>	\$ 31,410	\$ 367	\$ 112,108
Issuance of common stock for employee purchase		<u> </u>		<u>*</u>	Ψ 52,110	307	
plan	59,365	_	326	_	_	_	326
Exercise of stock options Issuance of common stock in settlement of restricted stock units, net of shares withheld for employee taxes, and stock	527,721	_	291	_	_	_	291
awards	43,042	_	(32)	_	_	_	(32)
expense Tax benefit from exercises of stock-based payment		_	4,236	-	_	_	4,236
awards	_	_	109		_	_	109
principle (see Note 1) Components of other comprehensive loss:	_	-		_	3,523	(3,523)	_
Net loss Other comprehensive income, net of full valuation allowance	-	_	_		(17,679)	_	(17,679)
on tax effect Comprehensive	_	_	_	_		1,494	1,494
loss Balance at December 31,							(16,185)
2009	13,436,163	<u>\$ 13</u>	\$ 85,248	<u>\$</u>	\$ 17,254	\$ (1,662)	\$ 100,853

The accompanying notes are an integral part of these consolidated financial statements.

CUTERA, INC. CONSOLIDATED STATEMENTS OF STOCKHOLDERS' EQUITY (continued)

Service Control									Accumulated							
3 (4.4) (4.4)	Common Stock			Lead of				ditional Paid-in		Deferred tock-Based	n.	etained	Cor	Other nprehensive	E4.	Total ockholders'
	Shares		ount	_	aıu-ın Capital		оск-ваseu mpensation		rnings		Income (loss)		Equity			
Balance at December 31,			ount		мриш				····							
2009	13,436,163	\$	13	\$	85,248	\$		\$	17,254	\$	(1,662)	\$	100,853			
Issuance of common stock												•				
for employee purchase	42.050				206								206			
plan	43,859		_		306				_		· —		306			
options	90,362		1		337		_						338			
Issuance of common stock	70,302		•		337											
in settlement of																
restricted stock units, net																
of shares withheld for			-													
employee taxes, and	50.220				(126)						• •		(126)			
stock awards Share-based compensation	59,329				(126)		_		_				(120)			
expense	_		_		4,650		_		_		´		4,650			
Tax benefit from exercises					.,											
of stock-based payment																
awards	_				8		_						. 8			
Components of other							,									
comprehensive loss:									(10,518)		<u> </u>		(10,518)			
Net loss Other comprehensive	_						_		(10,516)				(10,510)			
income, net of full																
valuation allowance																
on tax effect	·· —				. —		_		_		(94)		(94)			
Comprehensive					-								(10.610)			
loss	<u> </u>		=	_				-		_			(10,612)			
Balance at December 31,	12 620 712	ø	14.	ď	00.422	\$		æ	6,736	œ.	(1,756)	œ.	95,417			
2010	13,629,713	<u> </u>	14	<u> </u>	90,423	Φ.		<u> </u>	0,730	Φ	(1,730)	Ψ_	73,417			

The accompanying notes are an integral part of these consolidated financial statements.

CUTERA, INC. CONSOLIDATED STATEMENTS OF CASH FLOWS (in thousands)

	Year 1	r 31.	
	2010	2009	2008
Cash flows from operating activities:			
Net loss	\$ (10,518)	\$ (17,679)	\$ (2,869)
Adjustments to reconcile net loss to net cash provided by (used in)	ere jak		
operating activities:		٠,	•
Stock-based compensation	4,650	4,236	5,220
Tax benefit (deficit) from stock-based compensation	8	109	(231)
Excess tax benefit related to stock-based compensation	(8)	(23)	(44)
Depreciation and amortization	717	860	904
Provision for excess and obsolete inventories	235	611	409
Other-than-temporary impairments of long-term investments		· . —	3,554
Change in allowance for doubtful accounts	(122)	525	52
Change in deferred tax asset net of valuation allowance	(116)	10,512	(1,892)
Gain on sale of marketable and long term investments, net	(74)	(103)	(6)
Other			6
Changes in assets and liabilities:			
Accounts receivable	(759)	1,940	4,848
Inventories	(275)	2,908	(2,803)
Other current assets	2,314	1,014	1,354
Accounts payable	215	(609)	(660)
Accrued liabilities	(2,646)	42	(4,739)
Deferred rent	(200)	(62)	74
Deferred revenue	(1,208)	(3,537)	1,101
Income tax liability	(272)	(703)	62
Net cash provided by (used in) operating activities	(8,059)	41	4,340
Cash flows from investing activities:			
Acquisition of property and equipment	(275)	(154)	(703)
Proceeds from sales of marketable and long-term investments	42,830	27,914	55,104
Proceeds from maturities of marketable investments	42,505	11,535	30,065
Purchase of marketable and long-term investments	(87,837)	(53,655)	(63,822)
Net cash provided by (used in) investing activities	(2,777)	(14,360)	20,644
Cash flows from financing activities:			
Proceeds from exercise of stock options and employee stock purchase	i= .		
plan	518	585	458
Excess tax benefit related to stock-based compensation	8	23	44
Net cash provided by financing activities	526	608	502
Net increase (decrease) in cash and cash equivalents	(10,310)	(13,711)	25,486
Cash and cash equivalents at beginning of year	22,829	36,540	11,054
Cash and cash equivalents at end of year.	\$ 12,519	\$ 22,829	\$ 36,540
Supplemental and non-cash disclosure of cash flow information:	Ψ 12,517	<u> </u>	<u> </u>
Cash paid (received) for income taxes	\$ 272	\$ (578)	\$ 2,098
Cause para (10001100) for micomo macs	Ψ 2/2	<u>ψ (3/6)</u>	Ψ 2,090

CUTERA, INC. NOTES TO CONSOLIDATED FINANCIAL STATEMENTS

NOTE 1—SUMMARY OF SIGNIFICANT ACCOUNTING POLICIES

Description of Operations and Principles of Consolidation.

Cutera, Inc. (Cutera or the Company) is a global provider of laser and light-based aesthetic systems for practitioners worldwide. The Company designs, develops, manufactures, and markets the CoolGlide, Xeo, Solera, GenesisPlus and Excel V (introduced in 2011) product platforms for use by physicians and other qualified practitioners to allow its customers to offer safe and effective aesthetic treatments to their customers. The Xeo and Solera platforms offer multiple hand pieces and applications, which allow customers to upgrade their systems (Upgrade revenue). In addition to systems and upgrade revenue, the Company generates revenue from the sale of post warranty service contracts, providing services for products that are out of warranty, Titan hand piece refills, and dermal fillers and cosmeceuticals.

Headquartered in Brisbane, California, the Company has wholly-owned subsidiaries in Australia, Canada, France, Japan, Spain, Switzerland and United Kingdom that market, sell and service its products outside of the United States. The Consolidated Financial Statements include the accounts of the Company and its subsidiaries. All inter-company transactions and balances have been eliminated.

Use of Estimates.

The preparation of Consolidated Financial Statements in conformity with generally accepted accounting principles in the United States of America (GAAP) requires the Company's management to make estimates and assumptions that affect the amounts reported and disclosed in the financial statements and the accompanying notes. Actual results could differ materially from those estimates. On an ongoing basis, the Company evaluates their estimates, including those related to warranty obligation, sales commission, accounts receivable and sales allowances, fair values of long-term investments, fair values of acquired intangible assets, useful lives of intangible assets and property and equipment, fair values of options to purchase the Company's common stock, recoverability of deferred tax assets, and effective income tax rates, among others. Management bases their estimates on historical experience and on various other assumptions that are believed to be reasonable, the results of which form the basis for making judgments about the carrying values of assets and liabilities.

Cash, Cash Equivalents, Marketable Investments, and Long-Term Investments.

The Company invests its cash primarily in money market funds and in highly liquid debt instruments of U.S. federal and municipal governments and their agencies. All highly liquid investments with stated maturities of three months or less from date of purchase are classified as cash equivalents; all highly liquid investments with stated maturities of greater than three months are classified as marketable investments. The majority of the Company's cash and investments are held in U.S. banks and its foreign subsidiaries maintain a limited amount of cash in their local banks to cover their short term operating expenses.

The Company determines the appropriate classification of its investments in marketable securities at the time of purchase and re-evaluates such designation at each balance sheet date. The Company's marketable securities have been classified and accounted for as available-for-sale. The Company may, or may not, hold securities with stated maturities greater than 12 months until maturity. In response to changes in the availability of and the yield on alternative investments as well as liquidity requirements, it occasionally sells these securities prior to their stated maturities. As these securities are viewed by the Company as available to support current operations, based on the provisions of the Financial Accounting Standards Board Accounting Standards Codification (ASC) topic 210, subtopic 10, securities with maturities beyond 12 months (such as variable rate demand notes) are classified as current assets under the caption marketable investments in the accompanying Consolidated Balance Sheets. These securities are carried at fair value, with the unrealized gains and losses reported as a component of stockholders' equity. Any realized gains or losses on the sale of marketable securities are determined on a specific identification method, and such gains and losses are reflected as a component of interest and other income, net.

The Company holds a variety of interest bearing auction rate securities (ARS) that represent investments in pools of student loan assets issued by the Federal Family Education Loan Program (FELP). At the time of acquisition, the majority of ARS investments were intended to provide liquidity via an auction process that resets the applicable interest rate at predetermined calendar intervals, allowing investors to either roll over their holdings or gain immediate liquidity by selling such interests at par. Since February 2008, uncertainties in the credit markets affected the majority of ARS investments and auctions for the Company's investments in these securities have failed to settle on their respective settlement dates. However, since 2009 \$5.1 million of ARS were redeemed at full par value. Maturity dates for the ARS investments in the Company's portfolio range from 2028 to 2043.

As of December 31, 2010, the Company had \$6.8 million of ARS classified as long-term investments. The Company has classified its ARS investment balance as long-term investments in the accompanying Consolidated Balance Sheet because of the Company's belief that it could take more than one year before they are readily marketable. The Company's ARS have been classified and accounted for as available-for-sale. These securities are carried at fair value with the unrealized gains and losses reported as a component of stockholders' equity. The estimated fair value of the Company's ARS investments was \$6.8 million at December 31, 2010 and \$7.4 million at December 31, 2009.

Fair Value Measurements.

Fair value is defined as the price that would be received to sell an asset or paid to transfer a liability in an orderly transaction between market participants at the measurement date In determining fair value, the Company utilizes valuation techniques that maximize the use of observable inputs and minimize the use of unobservable inputs to the extent possible as well as considers counterparty credit risk in its assessment of fair value. Carrying amounts of the Company's financial instruments, including cash equivalents, marketable investments, accounts receivable, accounts payable and accrued liabilities, approximate their fair values as of the balance sheet dates because of their generally short maturities.

The fair value hierarchy distinguishes between (1) market participant assumptions developed based on market data obtained from independent sources (observable inputs) and (2) an entity's own assumptions about market participant assumptions developed based on the best information available in the circumstances (unobservable inputs). The fair value hierarchy consists of three broad levels, which gives the highest priority to unadjusted quoted prices in active markets for identical assets or liabilities (Level 1) and the lowest priority to unobservable inputs (Level 3). The three levels of the fair value hierarchy are described below:

- Level 1: Quoted prices (unadjusted) in active markets that are accessible at the measurement date for assets or liabilities.
- Level 2: Directly or indirectly observable inputs as of the reporting date through correlation with market data, including quoted prices for similar assets and liabilities in active markets and quoted prices in markets that are not active. Level 2 also includes assets and liabilities that are valued using models or other pricing methodologies that do not require significant judgment since the input assumptions used in the models, such as interest rates and volatility factors, are corroborated by readily observable data from actively quoted markets for substantially the full term of the financial instrument.
- Level 3: Unobservable inputs that are supported by little or no market activity and reflect the use of significant management judgment. These values are generally determined using pricing models for which the assumptions utilize management's estimates of market participant assumptions.

Impairment of Marketable Investments and ARS Securities.

The Company reviews its marketable and long term investments for impairment on a quarterly basis. If it concludes that any of these investments are impaired, it determines whether such impairment is other-than-temporary. Factors that the Company considers to make such determination include the duration and severity of the impairment, the reason for the decline in value and the potential recovery period, and its intent to sell, or whether it is more likely than not that it will be required to sell, the investment before recovery.

Beginning April 1, 2009, if an entity intends to sell, or if it is more likely than not that we will be required to sell, an impaired debt security prior to recovery of its cost basis, the security is other-than-temporarily impaired and the full amount of the impairment is required to be recognized as a loss through earnings. Otherwise, losses on securities which are other-than-temporarily impaired are separated into:

- (iii) the portion of loss which represents the credit loss; or
- (iv) the portion which is due to other factors.

The credit loss portion is recognized as a loss through earnings while the loss due to other factors is recognized in other comprehensive income (loss), net of taxes and related amortization. At December 31, 2010, the Company had approximately \$8.3 million of par value ARS investments on which with it had recognized approximately \$1.5 million in unrealized other-than-temporary losses. Given the Company believed that such losses were not credit related, it has included them in accumulated comprehensive loss.

Concentration of Credit Risk and Other Risks and Uncertainties.

Financial instruments that potentially subject the Company to concentrations of risk consist principally of cash, cash equivalents, marketable investments and accounts receivable. The Company's cash and cash equivalents are primarily invested in deposits and money market accounts with two major banks in the United States. In addition, the Company has operating cash balances in banks in each of the international locations in which it operates. Deposits in these banks may exceed the amount of insurance provided on such deposits, if any. Management believes that these financial institutions are financially sound and, accordingly, believes that minimal credit risk exists. The Company has not experienced any losses on its deposits of cash and cash equivalents. Accounts receivable are typically unsecured and are derived from revenue earned from worldwide customers. The Company performs credit evaluations of its customers and maintains reserves for potential credit losses. Concentrations of accounts receivable balances are presented in Note 3 and segment, geographic and major customer information is presented in Note 10.

The Company invests in debt instruments—including bonds and ARS—of the U.S. Government, its agencies and municipalities. In addition, starting from 2010, the Company has invested in other high grade investments such as commercial paper and corporate bonds. By policy, the Company restricts its exposure to any single issuer by imposing concentration limits. To minimize the exposure due to adverse shifts in interest rates, the Company maintains investments at an average maturity (interest reset date for auction-rate securities and variable rate demand notes) of generally less than eighteen months.

The Company is subject to risks common to companies in the medical device industry, including, but not limited to, new technology innovations, dependence on key personnel, dependence on key suppliers, protection of proprietary technology, product liability and compliance with government regulations. To continue profitable operations, the Company must continue to successfully design, develop, acquire, manufacture and market its products. There can be no assurance that current or recently acquired products will continue to be accepted in the marketplace. Nor can there be any assurance that any future products can be developed or manufactured at an acceptable cost and with appropriate performance characteristics, or that such products will be successfully marketed, if at all. These factors could have a material adverse effect on the Company's future financial results and cash flows.

Future products developed or acquired by the Company may require additional approvals from the Food and Drug Administration or international regulatory agencies prior to commercial sales. There can be no assurance that the Company's products will continue to meet the necessary regulatory requirements. If the Company was denied such approvals or such approvals were delayed, it may have a materially adverse impact on the Company.

Inventories.

Inventories are stated at the lower of cost or market, cost being determined on a standard cost basis (which approximates actual cost on a first-in, first-out basis) and market being determined as the lower of replacement cost or net realizable value.

The Company includes demonstration units within inventories. Demonstration units are carried at cost and amortized over their estimated economic life of two years. Amortization expense related to demonstration units is recorded in cost of revenue or in the respective operating expense line based on which function and purpose it is being used for.

Proceeds from the sale of demonstration units are recorded as revenue and all costs incurred to refurbish the systems prior to sale are charged to cost of revenue.

Property and Equipment.

Property and equipment are stated at cost and depreciated on a straight-line basis over the estimated useful lives of the related assets, which is generally three years. Amortization of leasehold improvements is computed using the straight-line method over the shorter of the remaining lease term or the estimated useful life of the related assets. Upon sale or retirement of assets, the costs and related accumulated depreciation and amortization are removed from the balance sheet and the resulting gain or loss is reflected in operating expenses. Maintenance and repairs are charged to operations as incurred.

Intangible Assets.

Purchased technology sublicense and other intangible assets are presented at cost, net of accumulated amortization. The technology licenses are being amortized on a straight-line basis over their expected useful life of 9-10 years. and the other intangibles are being amortized over their expected useful life of two years.

Impairment of Long-lived Assets.

The Company reviews long-lived assets, including property and equipment, and intangible assets, for impairment whenever events or changes in business circumstances indicate that the carrying amount of the assets may not be fully recoverable. The Company would recognize an impairment loss when estimated undiscounted future cash flows expected to result from the use of the asset and its eventual disposition is less than its carrying amount. Impairment, if any, is measured as the amount by which the carrying amount of a long-lived asset exceeds its fair value. Through December 31, 2010, there have been no such impairments.

Warranty Obligations.

The Company historically provided a standard one-year or two-year warranty coverage on its systems. Beginning in September 2009, the Company changed its warranty policy to a one-year standard warranty on all systems. Warranty coverage provided is for labor and parts necessary to repair the systems during the warranty period. The Company accounts for the estimated warranty cost of the standard warranty coverage as a charge to costs of revenue when revenue is recognized. The estimated warranty cost is based on historical product performance. To determine the estimated warranty reserve, the Company utilizes actual service records to calculate the average service expense per system and applies this to the equivalent number of units exposed under warranty. The Company updates these estimated charges every quarter.

Revenue Recognition.

Product, Upgrade, Titan hand piece refill, and Dermal filler and cosmeceutical revenue is recognized when title and risk of ownership has been transferred, provided that:

- Persuasive evidence of an arrangement exists;
- The price is fixed or determinable;
- Delivery has occurred or services have been rendered; and
- Collectability is reasonably assured.

Transfer of title and risk of ownership occurs when the product is shipped to the customer or when the customer receives the product, depending on the nature of the arrangement. Revenue is recorded net of customer and distributor discounts. For sales transactions when collectability is not reasonably assured, the Company recognizes revenue upon receipt of cash payment. Sales to customers and distributors do not include any return or exchange rights. In addition the Company's distributor agreements obligate the distributor to pay the Company for the sale regardless of whether the distributor is able to resell the product. Shipping and handling charges are invoiced to customers based on the amount of products sold. Shipping and handling fees are recorded as revenue and the related expense as a component of cost of revenue.

The Company also offers customers extended service contracts. Revenue under service contracts is recognized on a straight-line basis over the period of the applicable service contract. Service revenue, from customers whose systems are not under a service contact, is recognized as the services are provided. Service revenue for the years ended December 31, 2010, 2009, and 2008 was \$13.2 million, \$13.2 million, and \$11.4 million, respectively.

Shipping and Handling Costs.

Amounts charged to customers and costs incurred by the Company related to shipping and handling are included in net sales and cost of goods sold, respectively.

Research and Development Expenditures.

Costs related to research, design, development and testing of products are charged to research and development expense as incurred. Expenses incurred primarily relate to employees, facilities, material, third party contractors and clinical and regulatory fees.

Advertising Costs.

Advertising costs are included as part of sales and marketing expense and are expensed as incurred. Advertising expenses were \$947,000 in 2010, \$891,000 in 2009, and \$1.9 million in 2008.

Stock-based Compensation.

The Company elected to use the Black-Scholes-Merton (BSM) pricing model to determine the fair value of stock options on the dates of grant. Restricted stock units (RSUs) and stock awards are measured based on the fair market values of the underlying stock on the dates of grant. Shares are issued on the vesting dates, net of the statutory withholding requirements to be paid by the Company on behalf of its employees. As a result, the actual number of shares issued will be fewer than the actual number of RSUs outstanding. Furthermore, the Company records the liability for withholding amounts to be paid by us as a reduction to additional paid-in capital when the shares are issued. Also, the Company recognizes stock-based compensation using the straight-line method.

The Company includes as part of cash flows from financing activities the benefits of tax deductions in excess of the tax-effected compensation of the related stock-based awards for options exercised and RSUs vested during the period. The amount of cash received from the exercise of stock options and employee stock purchases was \$518,000 in 2010, \$585,000 in 2009, and \$458,000 in 2008, and the total direct tax benefit (deficit) realized, including the excess tax benefit (deficit), from stock-based award activity was \$8,000 in 2010, \$109,000 in 2009, and (\$231,000) in 2008. The Company elected to account for the indirect effects of stock-based awards—primarily the research and development tax credit—through the Statement of Operations.

Income Taxes.

The Company recognizes income taxes under the liability method. The Company recognizes deferred income taxes for differences between the financial reporting and tax bases of assets and liabilities at enacted statutory tax rates in effect for the years in which differences are expected to reverse. The Company recognizes the effect on deferred taxes of a change in tax rates in income in the period that includes the enactment date. The Company has determined that its future taxable income will be sufficient to recover all of the deferred tax assets. However, should there be a change in their ability to recover the deferred tax assets, the Company could be required to record a valuation allowance against its deferred tax assets. This would result in an increase to the Company's tax provision in the period in which they determined that the recovery was not probable.

The measurement of deferred taxes often involves an exercise of judgment related to the computation and realization of tax basis. The deferred tax assets and liabilities reflect management's assessment that tax positions taken, and the resulting tax basis, are more likely than not to be sustained if they are audited by taxing authorities. Also, assessing tax rates that the Company expects to apply and determining the years when the temporary differences are expected to affect taxable income requires judgment about the future apportionment of our income among the states in which the Company operates. These matters, and others, involve the exercise of significant judgment. Any changes in our

practices or judgments involved in the measurement of deferred tax assets and liabilities could materially impact our financial condition or results of operations.

Valuation allowances are established when necessary to reduce deferred income tax assets to amounts that the Company believes are more likely than not to be recovered. The Company evaluates its deferred tax assets quarterly to determine whether adjustments to our valuation allowance are appropriate. In making this evaluation, the Company relies on its recent history of pre-tax earnings, estimated timing of future deductions and benefits represented by the deferred tax assets, and its forecasts of future earnings, the latter two of which involve the exercise of significant judgment. As of September 30, 2009, the Company could not sustain a conclusion that it was more likely than not that the Company would realize any of its deferred tax assets resulting from its cumulative losses reported in the recent past as well as other factors. Consequently, the Company established a valuation allowance against those deferred tax assets. The Company also performed this evaluation as of December 31, 2010, and determined the full valuation allowance was still required.

The Company establishes reserves for uncertain tax positions in accordance with the Income Taxes subtopic of the ASC. The subtopic prescribes the minimum recognition threshold a tax position is required to meet before being recognized in the financial statements. Additionally, the subtopic provides guidance on derecognition, measurement, classification, interest and penalties, and transition of uncertain tax positions. The impact of an uncertain income tax position on income tax expense must be recognized at the largest amount that is more-likely-than-not to be sustained. An uncertain income tax position will not be recognized if it has less than a 50% likelihood of being sustained. The Company has provided taxes and related interest and penalties due for potential adjustments that may result from examinations of open U.S. Federal, state and foreign tax years. If the Company ultimately determines that payment of these amounts are not more-likely-than-not, the Company will reverse the liability and recognize a tax benefit during the period in which the Company makes the determination. The Company will record an additional charge in the Company's provision for taxes in the period in which the Company determines that the recorded tax liability is less than the Company expects the ultimate assessment to be.

Comprehensive Income (loss).

Comprehensive income (loss) generally represents all changes in stockholders' equity except those resulting from investments or contributions by stockholders. The Company's unrealized gains and losses on marketable investments represent the only component of other comprehensive income that is excluded from net income (loss).

On April 1, 2009, the Company adopted updates issued by the Financial Accounting Standards Board (FASB) to the recognition and presentation of other-than-temporary impairments. A cumulative effect adjustment was required to accumulated earnings and a corresponding adjustment to accumulated other comprehensive income (loss) to reclassify the non-credit portion of previously other-than-temporarily impaired securities which were held at the beginning of the period of adoption and for which the Company does not intend to sell and it is more likely than not that the Company will not be required to sell such securities before recovery of the amortized cost basis. As a result of the implementation of this pronouncement, the Company reclassified the cumulative effect of the non-credit portion of previously recognized other-than-temporarily impaired adjustments of \$3.5 million by increasing accumulated earnings and decreasing accumulated other comprehensive loss.

Foreign Currency.

The U.S. dollar is the functional currency of the Company's subsidiaries. Monetary and non-monetary assets and liabilities are remeasured into U.S. dollars at period end and historical exchange rates, respectively. Sales and operating expenses are remeasured at average exchange rates in effect during each period, except for those expenses related to non-monetary assets which are remeasured at historical exchange rates. Gains or losses resulting from foreign currency transactions are included in net income (loss) and are insignificant for each of the three years ended December 31, 2010. The effect of exchange rate changes on cash and cash equivalents was insignificant for each of the three years presented in the period ended December 31, 2010.

New Accounting Standards.

In October 2009, the FASB amended the accounting standards for revenue recognition to remove tangible products containing software components and non-software components that function together to deliver the product's essential functionality from the scope of the software revenue recognition guidance. The software revenue recognition guidance was issued to address factors that entities should consider when determining whether the software and non-software components of a product function together to deliver the product's essential functionality. The software revenue recognition updates to the Codification will allow revenue arrangements in which software and non-software components deliver together a product's essential functionality to follow the multiple-deliverable revenue recognition criteria as opposed to the criteria applicable to software revenue recognition.

In addition, the FASB also amended the accounting standards for multiple deliverable revenue arrangements to:

- provide updated guidance on whether multiple deliverables exist, how the deliverables in an arrangement should be separated, and how the consideration should be allocated;
- require an entity to allocate revenue in an arrangement using estimated selling price (ESP) of deliverables if a vendor does not first have vendor-specific objective evidence (VSOE) of selling price or secondly does not have third-party evidence (TPE) of selling price; and
- eliminate the use of the residual method and require an entity to allocate revenue using the relative selling price method.

Multiple-element arrangements- A multiple-element arrangement includes the sale of one or more tangible product offerings with one or more associated services offerings, each of which are individually considered separate units of accounting. The determination of the Company's units of accounting did not change with the adoption of the new revenue recognition guidance, and as such the Company allocates revenue to each element in a multiple-element arrangement based upon the relative selling price of each deliverable. When applying the relative selling price method, the Company determines the selling price for each deliverable using VSOE of selling price, if it exists, or TPE of selling price. If neither VSOE nor TPE of selling price exist for a deliverable, the Company uses its best estimate of selling price for that deliverable. Revenue allocated to each element is then recognized when the other revenue recognition criteria are met for each element.

Both the above mentioned updates are effective for the Company from January 1, 2011 and the Company has elected to apply them prospectively to new or materially modified revenue arrangements after its effective date. The Company does not expect the adoption of this guidance to have a material impact on its Consolidated Financial Statements and results of operations.

NOTE 2—INVESTMENT SECURITIES:

The following tables summarize cash, cash equivalents, marketable securities and long term investments (in millions):

	Decem	ber 3	1,	
	2010		2009	
Cash and cash equivalents:	 			
Cash	\$ 1,989	\$	3,483	
Cash equivalents:	ŕ		1.75	
Money market funds	8,330		19,346	
Commercial paper	2,200		, -	
Total cash and cash equivalents	 12,519		22,829	
Marketable securities:		5		
U.S. government notes	2.070			
U.S. government agencies	24,087		_	
Municipal securities	15,011		76,680	
Commercial Paper	11,465		_	
Corporate debt securities	24,851			
ARS	_		100	
Total marketable securities	77,484		76,780	
Long-term investments in ARS	 6,784	_	7,275	
investments	\$ 96,787	\$	106,884	

The following table summarizes unrealized gains and losses related to our marketable investments and long term investments, both designated as available-for-sale (in thousands):

		*,																
	Amortized Cost		Amortized Cost										Unre	oss alized iins	Gross Unrealized Losses			Fair . Market Value
Cash and cash equivalents	\$	12,519	\$		\$		\$	12,519										
Marketable investments																		
U.S. government notes		2,069		1				2,070										
U.S. government agencies		24,088		17		(18)		24,087										
Municipal securities		15,029		2		(20)		15,011										
Commercial Paper		11,459		7		(1)		11,465										
Corporate debt securities		24,825		55		(29)		24,851										
Total marketable securities	_	77,470		82		(68)		77,484										
Long-term investments in ARS		8,325				(1,541)		6,784										
Total cash, cash equivalents, marketable securities	ø	09 214	ф	02	er.	(1.600)	e	06 797										
and long term investments	<u>\$</u>	98,314	\$	82	<u>\$</u>	(1,609)	\$	96,787										
December 31, 2009	.	mortized , Cost	Unr	ross ealized ains	Uı	Gross nrealized Losses		Fair Market Value										
Cash and cash equivalents	\$	22,829	\$		\$		\$	22,829										
Marketable investments:		,																
Municipal securities		76,512		182		(14)		76,680										
ARS		100		_				100										
Total marketable investments		76,612		182		(14)		76,780										
Long-term investments in ARS		8,875				(1,600)		7,275										
Total cash, cash equivalents, marketable securities	-																	
and long term investments	\$	108,316	\$	182	\$	(1,614)	\$	106,884										

The following table summarizes the estimated fair value of our marketable investments and long term investments classified by the contractual maturity date of the security as of December 31, 2010 (in thousands):

	 Mount
Due in less than one year (fiscal year 2011)	\$ 41,796
Due in 1 to 3 years (fiscal year 2012- 2013)	35,688
Due in 3 to 5 years (fiscal year 2014-2015)	_
Due in 5 to 10 years (fiscal year 2016-2021)	
Due in greater than 10 years (fiscal year 2022 and beyond)	 6,784
	\$ 84,268

Fair Value Measurements

As of December 31, 2010, financial assets measured and recognized at fair value on a recurring basis and classified under the appropriate level of the fair value hierarchy as described above was as follows (in thousands):

	Level 1		Level 1 Level 2		Level 3			Total
Cash equivalents	\$	10,531	\$	_	\$		\$	10.531
Short term marketable investments:								. ,
Available-for-sale securities		. —		77,484				77,484
Long-term investments:				, -				, , , , , , ,
Available-for-sale ARS				_	ì	6,784	* ,	6,784
Total assets at fair value	\$	10,531	\$	77,484	\$	6,784	\$	94,799

The Company's Level 1 financial assets are money market funds and highly liquid debt instruments of U.S. federal and municipal governments and their agencies with stated maturities of three months or less from the date of purchase, whose fair values are based on quoted market prices. The Company's Level 2 financial assets are highly liquid debt instruments of U.S. federal and municipal governments and their agencies with stated maturities of greater than three months, whose fair values are obtained from readily-available pricing sources for the identical underlying security that may, or may not, be actively traded.

At December 31, 2010, observable market information was not available to determine the fair value of the Company's ARS investments. Therefore, the fair value is based on broker-provided valuation models that relied on Level 3 inputs including those that are based on expected cash flow streams and collateral values, assessments of counterparty credit quality, default risk underlying the security, market discount rates and overall capital market liquidity. The valuation of the Company's ARS investment portfolio is subject to uncertainties that are difficult to predict. Factors that may impact the valuations in the future include changes to credit ratings of the securities, as well as to the underlying assets supporting those securities, rates of default of the underlying assets, underlying collateral value, discount rates, counterparty risk and ongoing strength and quality of market credit and liquidity. These financial instruments are classified within Level 3 of the fair value hierarchy.

The table presented below summarizes the change in carrying value associated with Level 3 financial assets, which represents the Company's investment in ARS and classified as long-term investments, for the year ended December 31, 2010 (in thousands):

	Dece	mber 31	l, 2010
Balance at December 31, 2009	\$	•	7,275
Total gains or losses (realized or unrealized)			
Included in earnings (or changes in net assets)	.:		
Included in other comprehensive income (loss)			(26)
Purchases, issuance, and settlements			(465)
Transfers in and/or out of Level 3		•	_
Balance at December 31, 2010			

NOTE 3—BALANCE SHEET DETAIL:

Accounts Receivable:

Trade accounts receivable are recorded at the invoiced amount and do not bear interest. The allowance for doubtful accounts is the Company's best estimate of the amount of probable credit losses existing in accounts receivable and is based on historical write-off experience and any specific customer issues that have been identified. Account balances are charged off against the allowance when it is probable the receivable will not be recovered. The Company had one customer who accounted for 10% at December 31, 2010 and 29% at December 31, 2009 of the Company's total accounts receivable balance.

Inventories:

Inventories consist of the following (in thousands):

		,		
		2010		2009
Raw materials	\$	4,204	\$	3,775
Finished goods		2,243		2,633
Total	\$	6,447	\$	6,408

Property and Equipment, net:

Property and equipment, net consists of the following (in thousands):

	December 31,			
	2	2010		2009
Leasehold improvements	\$	361	\$	347
Office equipment and furniture		2,702		2,610
Machinery and equipment		2,688		2,519
		5,751	-	5,476
Less: Accumulated depreciation		(5,154)		(4,629)
Property and equipment, net	\$	597	\$	847

Depreciation expense related to property and equipment was \$525,000 in 2010, \$664,000 in 2009, and \$702,000 in 2008.

Intangible Assets:

Intangible assets were principally comprised of a patent sublicense acquired from Palomar in 2006, a technology sublicense acquired in 2002 and other intangible assets acquired in 2007. The components of intangible assets at December 31, 2010 and 2009 were as follows (in thousands):

	Gross Carrying Amount		Am	umulated ortization Amount	Net mount
December 31, 2010					
Patent sublicense	\$	1,218	\$	656	\$ 562
Technology sublicense		538		463	75
Total	\$	1,756	\$	1,119	\$ 637
December 31, 2009			·		
Patent sublicense	\$	1,218	\$	517	\$ 701
Technology sublicense		538		410	128
Other intangibles		20		20	·
Total	\$	1,776	\$	947	\$ 829

Amortization expense for intangible assets was \$192,000 in 2010, \$196,000 in 2009, and \$202,000 in 2008.

Based on intangible assets recorded at December 31, 2010, and assuming no subsequent additions to, or impairment of the underlying assets, the remaining estimated annual amortization expense is expected to be as follows (in thousands):

Year ending December 31,				Amc	ount
2011					192
2012		 	 		158
2013					138
2014	and the second s				138
2015					11
Total		 	 	 \$	637

Accrued Liabilities:

Accrued liabilities consist of the following (in thousands):

$\mathcal{F}_{i} = \mathcal{F}_{i}$	and the same of th			Decem	ber 31,	
				2010		2009
Payroll and related expenses	• • • • • • • • • • • • • • • • • • • •		. \$	3,035	\$	3,216
Sales tax				809		748
Warranty		• • • • • • • • • • • • • • • • • • • •		796		1,049
Royalty				475		476
Professional fees				335		733
Customer deposits				131		667
Sales and marketing accruals				131		325
Other	· · · · · · · · · · · · · · · · · · ·			482		884
Litigation accrual - Telephone Cor	nsumer Protection Act (s	see Note 11)	•	—		950
Total	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	. \$	6,194	\$	9,048

NOTE 4—WARRANTY AND SERVICE CONTRACTS:

The Company has a direct field service organization in the United States. Internationally, the Company provides direct service support through its wholly-owned subsidiaries in Australia, Canada, France, Japan, Spain and Switzerland as well as through a network of distributors and third-party service providers in several other countries where it does not have a direct presence. The Company provides a warranty with its products, depending on the type of product. After the original warranty period, maintenance and support are offered on a service contract basis or on a time and materials basis. The Company currently provides for the estimated cost to repair or replace products under warranty at the time of sale.

Warranty Accrual (in thousands):

	December 31,			
		2010		2009
Balance at beginning of year	\$	1,049	\$	1,916
Add: Accruals for warranties issued during the year		3,061		2,059
Less: Settlements made during the year		(3,314)		(2,926)
Balance at end of year	\$	796	\$	1,049

Deferred Service Contract Revenue (in thousands):

	· . · · · · · · · · · · · · · · · · · ·		1,		
	* **		2010		2009
Balance at beginning of year	*********	\$	8,128	\$	11,665
Add: Payments received			8,254		6,585
Less: Revenue recognized			(9,617)		(10,122)
Balance at end of year		\$	6,765	\$	8,128

Costs incurred under service contracts amounted to \$4.3 million in 2010, \$4.7 million in 2009, and \$4.4 million in 2008, and are recognized as incurred.

NOTE 5—STOCKHOLDERS' EQUITY, STOCK PLANS AND STOCK-BASED COMPENSATION EXPENSE:

Stock Option Plans.

As of December 31, 2010, the Company had the following stock-based employee compensation plans:

2004 Employee Stock Purchase Plan.

On January 12, 2004, the Board of Directors adopted the 2004 Employee Stock Purchase Plan. A total of 200,000 shares of common stock were reserved for issuance pursuant to the 2004 Employee Stock Purchase Plan. Under the 2004 Employee Stock Purchase Plan, or 2004 ESPP, eligible employees are permitted to purchase common stock at a discount through payroll deductions. The 2004 ESPP offering and purchase periods are for approximately six months. Shares of common stock eligible for purchase are increased on the first day of each fiscal year by an amount equal to the lesser of (i) 600,000 shares, (ii) 2.0% of the outstanding shares of common stock on such date or (iii) an amount as determined by the Board of Directors. The Company's Board of Directors voted not to increase the shares available for future grant on January 1, 2010 and reserved 256,121 shares on January 1, 2009. The price of the common stock purchased is the lower of 85% of the fair market value of the common stock at the beginning of an offering period or at the end of the offering period. Under the 2004 ESPP the Company issued 43,859 shares in 2010 and 59,365 shares in 2009. At December 31, 2010, 1,102,097 shares remained available for future issuance.

2004 Equity Incentive Plan and 1998 Stock Plan.

In 1998, the Company adopted the 1998 Stock Plan, or 1998 Plan, under which 4,650,000 shares of the Company's common stock have been reserved for issuance to employees, directors and consultants.

On January 12, 2004, the Board of Directors adopted the 2004 Equity Incentive Plan. A total of 1,750,000 shares of common stock were originally reserved for issuance pursuant to the 2004 Equity Incentive Plan. In addition, the shares reserved for issuance under the 2004 Equity Incentive Plan included shares reserved but un-issued under the 1998 Plan and shares returned to the 1998 Plan as the result of termination of options or the repurchase of shares.

Shares of common stock approved under the 2004 Equity Incentive Plan was increased on the first day of each fiscal year, commencing in 2005, by an amount equal to the lesser of: (i) 5% of the outstanding shares on the first day of such year; (b) 2 million shares; or, (c) an amount determined by the Board of Directors. The Company added 636,922 shares to the 2004 Equity Incentive Plan on January 1, 2009. During 2009, the 2004 Equity Incentive Plan was amended to remove this feature beginning in 2010.

Options granted under the 1998 Plan and 2004 Equity Incentive Plan may be incentive stock options or non-statutory stock options. Stock purchase rights may also be granted under the 2004 Equity Incentive Plan. Incentive stock options may only be granted to employees. The Board of Directors determines the period over which options become exercisable, however, except in the case of options granted to officers, directors and consultants, options shall become exercisable at a rate of no less than 20% per year over five years from the date the options are granted. Options are to be granted at an exercise price not less than the fair market value per share on the grant date for incentive options or 85% of fair market value for nonqualified stock options. For employees holding more than 10% of the voting rights of all classes of stock, the exercise price shall not be less than 110% of the fair market value per share on the grant date. Options granted under the Plan to employees generally become exercisable 25% on the first anniversary of the vesting commencement date and an additional 1/48th of the total number of shares subject to the option shares shall become exercisable on the last day of each calendar month thereafter until all of the shares have become exercisable. Unvested options that have been exercised are subject to repurchase upon termination of the holder's status as an employee, director or consultant. The contractual term of the options granted is either five, seven or ten years.

Under the 2004 Equity Incentive Plan, in May 2010 the Company's Board of Directors approved the grant of 37,266 Restricted Stock Units (RSU) to non-employee members of the Board of Directors that vested immediately on the date of grant. In addition, the Company's Board of Directors granted 109,025 RSUs to certain members of the Company's management that vested one-third on June 1, 2010 and one-third will vest on June 1, 2011 and 2012. The Company measured the fair market values of the underlying stock on the dates of grant and recognizes the share-based compensation expense using the straight-line method over the vesting period.

The Company issues new shares upon the exercise of options, restricted stock units and ESPP shares.

Option Exchange Program.

In July 2009, the Company completed its Option Exchange Program for its employees to exchange certain options outstanding for new options to purchase shares of the Company's common stock. As a result, options to purchase 864,373 shares of the Company's common stock were cancelled and new options to purchase up to 447,841 shares of the Company's common stock were issued in exchange. The new options have an exercise price per share of \$8.49, the closing price of the Company's common stock as reported on the Nasdaq Global Select Market on the date that the offer expired and Option Exchange Program was completed, are unvested as of the grant date, and subject to an additional six (6) months of vesting over and above the vesting schedule of the surrendered options.

Given the Option Exchange Program was designed to be approximately a "value-for-value" exchange, the Company did not incur any significant additional non-cash compensation charges as the fair value of the replacement options was approximately equal to or less than the fair value of the surrendered options. The Company determined the fair value of stock options using the Black Scholes valuation model.

Option Activity.

Activity under the 1998 Plan and 2004 Equity Incentive Plan is summarized as follows:

		· ·	Options	Outstanding	
	Shares Available For Grant	Number of Shares	Veighted- Average Exercise Price	Weighted- Average Remaining Contractual Life (in years)	Aggregate Intrinsic Value (in \$ millions) ⁽¹⁾
Balances as of December 31, 2007	2,047,649	2,417,575	\$ 14.22		(311 4 11111191119)
Additional shares reserved	636,922		_		
Options granted	(888,150)	888,150	\$ 10.77		4
Options exercised		(8,449)	\$ 5.39		
Options cancelled or forfeited	215,543	(215,543)	\$ 18.69		
Restricted stock units cancelled or forfeited	1,125				
Balances as of December 31, 2008	2,013,089	3,081,733	\$ 12.94	4.58	\$ 6.0
Options granted ⁽²⁾	(1,409,371)	1,409,371	\$ 8.51		
Options exercised	_	(527,721)	\$ 0.55		
Options cancelled (expired or forfeited) ⁽²⁾	1,270,828	(1,270,828)	\$ 17.55		
Stock awards granted	(36,540)			-	
Restricted stock units cancelled (expired or forfeited)	2,375	_			
Balances as of December 31, 2009	1,840,381	2,692,555	\$ 10.87	5.05	\$ 1.6
Options granted ⁽²⁾	(961,500)	961,500	\$ 10.14		
Options exercised		(90,362)	\$ 3.74		
Options cancelled (expired or		` , ,			
forfeited) ⁽²⁾	267,274	(267,274)	9.91		
Stock awards granted	(146,291)				
Restricted stock units cancelled (expired or					
forfeited)	5,583		_		
Balances as of December 31, 2010	1,005,447	3,296,419	\$ 10.93	4.4	\$ 1.1
Exercisable as of December 31, 2010		1,679,268	\$ 12.12	3.29	\$ 1.0

Based on the closing stock price of the Company's stock of \$8.29 on December 31, 2010, \$8.51 on December 31, 2009 and \$8.87 on December 31, 2008.

Included in options granted and options cancelled are shares granted and cancelled in connection with the Company's Option Exchange Program in 2009 (see 'Option Exchange Program' above for more details).

The aggregate intrinsic value in the table above represents the total pre-tax intrinsic value (the aggregate difference between the Company's closing stock price on the last trading day of the fiscal year 2010 and the exercise price, multiplied by the number of in-the-money options) that would have been received by the option holders had all option holders exercised their options on December 31, 2010. The aggregate intrinsic amount changes based on the fair market value of the Company's common stock. Total intrinsic value of options exercised was \$128,000 in 2010, \$3.2 million in 2009, and \$57,000 in 2008. The options outstanding and exercisable at December 31, 2010 were in the following exercise price ranges:

	Options	Outstanding	Option	s Exercisable	
Range of Exercise Prices	Number Outstanding	Weighted-Average Remaining Contractual Life (in years)	Number Outstanding	Weighted-Averag Exercise Price	e .
\$ 2.50-\$7.73	332,350	2.78	265,932	\$ 4.3	37
\$ 8.02–\$8.02	23,542	3.37	11,511	8.0	02
\$ 8.49–\$8.49	429,753	3.19	336,908	8.4	49
\$ 8.56–\$8.56	3,750	5.56	1,407	8.5	56
\$ 8.66–\$8.66	725,324	5.14	309,446	8.0	66
\$ 8.81–\$9.74	66,000	4.70	40,188	9.0	62
\$ 10.24-\$10.24	791,000	6.21	_	-	_
\$ 10.43–\$13.30	343,433	4.24	144,836	11.8	83
\$ 13.54–\$23.75	449,017	3.19	447,893	18.7	79
\$ 24.46–\$34.45	132,250	2.14	121,147	24.9	96
\$ 2.50–\$34.45	3,296,419	4.40	1,679,268	\$ 12.	12

As of December 31, 2009 there were 1,253,360 options that were exercisable at a weighted average exercise price of \$12.54.

Restricted Stock Units and Stock Awards.

Information with respect to restricted stock units activity is as follows (in thousands):

	Number of Shares	ighted-Average rant-Date Fair Value	Aggregate air Value ⁽¹⁾ n thousands)
Outstanding at December 31, 2009		\$ _	
Granted	146,291	\$ 10.09	
Vested ⁽²⁾	(73,612)	\$ 10.09	\$ 680 ⁽³⁾
Forfeited		\$ 10.09	
Outstanding at December 31, 2010	67,096	\$ 10.09	

(1) Represents the value of the Company's stock on the date that the restricted stock units vest.

The number of restricted stock units vested includes shares that the Company withheld on behalf of the employees to satisfy the statutory tax withholding requirements.

On the grant date, the fair value for these vested awards was \$743,000.

Stock-Based Compensation.

Stock-based compensation expense for stock options, restricted stock units, stock awards and ESPP shares for the year ended December 31, 2010, 2009 and 2008 was as follows (in thousands):

	Year Ended December 31,					
		2010		2009		2008
Stock options	\$	3,628	\$	3,763	\$	4,783
RSUs and Stock awards		927		360		257
ESPP		95		113		<u> 180</u>
Total share-based compensation expense		4,650		4,236		5,220
Tax effect on share-based compensation at the marginal tax rates		(1,725)		(1,452)		(1,788)
Net share-based compensation expense	\$	2,925	\$	2,784	<u>\$</u>	3,432

Total pre-tax stock-based compensation expense by department recognized during the year ended December 31, 2010, 2009 and 2008 was as follows (in thousands):

	Year Ended December 31,						
the state of the s		2010		2009		2008	
Cost of revenue	\$	724	\$	717	\$	846	
Sales and marketing		1,189		1,044		1,657	
Research and development		629		473		628	
General and administrative		2,108		2,002		2,089	
Total share-based compensation expense	\$	4,650	\$	4,236	\$	5,220	

As of December 31, 2010, the unrecognized compensation cost, net of expected forfeitures, was \$5.5 million for stock options and stock awards, which will be recognized using the straight- line attribution method over an estimated weighted-average remaining amortization period of 2.40 years. For ESPP, the unrecognized compensation cost, net of expected forfeitures, was \$29,000, which will be recognized using the straight- line attribution method over an estimated weighted-average amortization period 0.33 years.

Valuation Assumptions and Fair Value of Stock Option and ESPP Grants.

The Company uses the Black-Scholes option pricing model to estimate the fair value of options granted under its equity incentive plans and rights to acquire stock granted under its employee stock purchase plan. The Company based the weighted average estimated values of employee stock option grants and rights granted under the employee stock purchase plan, as well as the weighted average assumptions used in calculating these values, on estimates at the date of grant, as follows:

		Stock	k Options		Sto	ck Pu	rchase Pla	n	
	·	2010	2009	 2008	2010		2009		2008
Estimated fair value of grants during		24.							
the year	\$	3.76 \$	3.93	\$ 5.29	\$ 2.41	\$	2.39	\$	4.52
Expected term (in years) ⁽¹⁾		3.84	4.23	4.68	0.50		0.50		0.50
Risk-free interest rate ⁽²⁾		1.73%	2.6%	3.2%	0.2%	,	0.1%		1.9%
Volatility ⁽³⁾		46%	55%	55%	40%	,	52%		51%
Dividend yield ⁽⁴⁾		%	%	%	%	,	%		-%

- The expected term represents the period during which the Company's stock-based awards are expected to be outstanding. The estimated term is based on historical experience of similar awards, giving consideration to the contractual terms of the awards, vesting requirements, and expectation of future employee behavior, including post-vesting terminations. Prior to 2008, the Company used the simplified method of calculating expected life described in SAB 107, Share Based Payment, due to significant differences in the vesting and contractual life of current option grants compared to its historical grants, as well as limited data of historical exercise patterns since the Initial Public Offering (IPO) of its common stock.
- The risk-free interest rate is based on U.S. Treasury debt securities with maturities close to the expected term of the option as of the date of grant.
- Expected volatility is a 50%/50% blend of implied and historical volatility. The Company has determined that this is a more reflective measure of market conditions and a better indicator of expected volatility, than its limited historical volatility since the IPO of its common stock.
- The Company has not historically issued any dividends and does not expect to do so in the foreseeable future.

The Company periodically estimates forfeiture rates based on its historical experience within separate groups of employees and adjusts the share-based payment expense accordingly.

NOTE 6: COMMON STOCK REPURCHASES

Restricted Stock Unit Withholdings

The Company issues restricted stock units as part of its equity incentive plans, which are described more fully in "Note 5—Stockholders' Equity, Stock Plans and Stock-Based Compensation Expense." For the majority of restricted stock units granted, the number of shares issued on the date the restricted stock units vest is net of the statutory withholding requirements paid on behalf of the employees. The Company withheld 14,283 in 2010, 3,934 in 2009, and 4,992 in 2008, shares of common stock to satisfy its employees' tax obligations of \$126,000 in 2010, \$32,000 in 2009, and \$51,000 in 2008. The Company paid this amount in cash to the appropriate taxing authorities. Although shares withheld are not issued, they are treated as common stock repurchases for accounting and disclosure purposes, as they reduce the number of shares that would have been issued upon vesting.

NOTE 7—INCOME TAXES:

The Company files income tax returns in the U.S. federal and various state and local jurisdictions and foreign jurisdictions. The components of the provision for income taxes are as follows (in thousands):

	*	Ye	ar End	ed December 3	1,	
and the second of the second o		2010		2009		2008
Current:	• •					
Federal	\$	(154)	\$	(1,973)	\$	1,009
State		. 37		32		305
StateForeign		235		338		382
		118		(1,603)		1,696
Deferred:						
Federal		(45)		9,686		(2,313)
State		45		871		(78)
Foreign		(116)		. (46)		(97)
		(116)		10,511		(2,488)
Provision (benefit) for income taxes	\$	2	\$	8,908	\$	(792)

The Company's deferred tax asset consists of the following (in thousands):

		Deceml	ber 31,	
		2010		2009 ⁽¹⁾
Net operating loss ⁽¹⁾	\$.	6,281	\$	2,403
Stock-based compensation		5,644		4,499
Other accruals and reserves		3,385		4,631
Credits		1,488		1,184
Capital loss		558		539
Foreign		388		272
Accrued warranty		303		401
Other		÷63		78
Deferred tax asset	3.	. 18,110		14,007
Depreciation and amortization		146		103
Net deferred tax asset before valuation allowance	- :-	18,256		14,110
Valuation allowance ⁽¹⁾		(17,868)		(13,838)
Net deferred tax asset after valuation allowance	\$	388	\$	272

⁽¹⁾ The Company revised the 2009 tax footnote to reduce deferred tax assets by approximately \$1.1 million related to future tax benefits for net operating losses that were not properly recorded in the previous period. This reduction in deferred taxes was offset by a corresponding reduction in the valuation allowance, and as such had no impact to the Consolidated Financial Statements, earnings per share, statement of cash flows, or statement of equity for any period presented.

Due to uncertainties surrounding the realization of U.S. deferred tax assets through future taxable income, the Company has provided a full valuation allowance and, therefore, has not recognized any benefits from the U.S. net operating losses and other U.S. deferred tax assets. The valuation allowance balance was approximately \$13.8 million, \$1.4 million and \$0 at January 1, 2010, 2009 and 2008, respectively. Additions to the valuation allowance were approximately \$5.4 million, \$13.1 million and \$1.4 million for the years ended December 31, 2010, 2009 and 2008, respectively. Deductions to the valuation allowance were approximately \$1.4 million, \$660,000 and \$0 for the years ended December 31, 2010, 2009 and 2008, respectively. The valuation allowance balance was approximately \$17.9 million, \$13.8 million and \$1.4 million at December 31, 2010, 2009 and 2008, respectively.

The differences between the U.S. federal statutory income tax rate to the Company's effective tax are as follows:

	Year	Ended December 31,	
TIO C.1. 1	2010	2009(1)	2008
U.S. federal statutory income tax rate	35.00%	35.00%	35.00%
State tax rate, net of federal benefit	2.81	(0.86)	(2.38)
Benefit for research and development credit	2.97	1.06	11.07
Decreases related to release of unrecognized tax benefits		1.00	11.07
due to the lapsing of statute of limitations	2.59	0.71	2.66
Tax-exempt interest	0.98	5.42	27.85
Meals and entertainment	(0.63)	(0.76)	(3.45)
Income tax refund	(1.13)	11.00	(3.43)
Stock-based compensation	(1.54)	(8.91)	(5 (5)
Valuation allowance ⁽¹⁾	, ,	` ,	(5.65)
Other	(38.31)	(142.18)	(37.34)
Effective toy rete	(2.74)	(2.05)	(6.13)
Effective tax rate	0.00%	(101.57)%	21.63%

⁽¹⁾ The Company revised the 2009 tax footnote to reduce deferred tax assets by approximately \$1.1 million related to future tax benefits for net operating losses that were not properly recorded in the previous period. This reduction in deferred taxes was offset by a corresponding reduction in the valuation allowance, and as such had no impact to the Consolidated Financial Statements, earnings per share, statement of cash flows, or statement of equity for any period presented.

The Company recognizes deferred tax assets for the expected future tax consequences of temporary differences between the financial reporting and tax bases of assets and liabilities, and for operating losses and tax credit carryforwards. The Company records a valuation allowance to reduce the deferred tax assets to their estimated realizable value, when it is more likely than not that it will not be able to generate sufficient future taxable income to realize the net carrying value. The Company reviews the deferred tax asset and valuation allowance on a quarterly basis, and considers whether positive and negative evidence exists to effect the realization of deferred tax assets. After considering both the positive and negative evidence as of September 30, 2009, the Company determined that it was not more-likely-than-not that it would realize the full value of its deferred tax assets. As a result, in 2009 the Company established a valuation allowance of \$10.2 million against the net deferred tax asset balance as of December 31, 2008. In addition, the Company recorded a valuation allowance against its deferred tax assets generated in 2009 and 2010, which resulted in a valuation allowance of \$17.9 million as of December 31, 2010.

As of December 31, 2010, the Company had cumulative net operating loss carry-forwards for federal and state income tax reporting purposes of approximately \$17.0 million and \$5.6 million, respectively. The federal net operating loss carry-forwards expire through the year 2030 and the state net operating loss carry-forwards expire at various dates through the year 2030. Such net operating losses consist of excess tax benefits from employee stock option exercises and have not been recorded in the Company's deferred tax assets.

As of December 31, 2010, the Company had research and development tax credits for federal and state income tax purposes of approximately \$2.9 million and \$3.2 million, respectively. The federal research and development tax credits expire through the year 2030. The state research and development credits can be carried forward indefinitely, except for \$284,000, which will expire at various dates through the year 2020. The Company maintained a valuation allowance against these tax credits at December 31, 2010.

At December 31, 2010, the Company had an aggregate of approximately \$3 million of unremitted earnings of foreign subsidiaries that have been, or are intended to be, indefinitely reinvested for continued use in foreign operations. If the total undistributed earnings of foreign subsidiaries were remitted, a significant amount of the additional tax would be offset by the allowable foreign tax credits. The Company has determined that it is not practical for it to determine the additional tax of remitting these earnings.

Uncertain Tax Positions

The Company establishes reserves for uncertain tax positions in accordance with the ASC. The subtopic prescribes the minimum recognition threshold a tax position is required to meet before being recognized in the financial statements. Additionally, the subtopic provides guidance on derecognition, measurement, classification, interest and penalties, and transition of uncertain tax positions. The impact of an uncertain income tax position on income tax expense must be recognized at the largest amount that is more-likely-than-not to be sustained. An uncertain income tax position will not be recognized if it has less than a 50% likelihood of being sustained. The Company has provided taxes and related interest and penalties due for potential adjustments that may result from examinations of open U.S. Federal, state and foreign tax years. If the Company ultimately determines that payment of these amounts are not more-likely-than-not, the Company will reverse the liability and recognize a tax benefit during the period in which the Company makes the determination. The Company will record an additional charge in the Company's provision for taxes in the period in which the Company determines that the recorded tax liability is less than the Company expects the ultimate assessment to be. The Company's policy to include interest and penalties related to gross unrecognized tax benefits within the provision for income taxes did not change.

The following table summarizes the activity related to the Company's gross unrecognized tax benefits from December 31, 2009 to December 31, 2010 (in thousands):

	Year Ended December 31,							
		2010		2009		2008		
Balance at beginning of year	\$	787	\$	1,640	\$	1,500		
Increases related to prior year tax positions				88		· · · · —		
Decreases related to prior year tax positions		(29)		(857)		(98)		
Increases related to current year tax positions	•	24		29		258		
Decreases related to lapsing of statute of limitations		(227)		(113)		(20)		
Balance at end of year	\$	555	\$	787	\$	1,640		

The Company's total unrecognized tax benefits that, if recognized, would affect its effective tax rate were approximately \$405,000 and \$737,000 as of December 31, 2010 and 2009, respectively. The Company had accrued approximately \$71,000 and \$117,000 for payment of interest as of December 31, 2010 and 2009, respectively. Interest included in the provision for income taxes was not significant in all the periods presented. The Company has not accrued any penalties related to its uncertain tax positions as it believes that it is more likely than not that there will not be any assessment of penalties. The Company expects that the amount of unrecognized tax benefits will not change within the next 12 months.

NOTE 8—NET LOSS PER SHARE:

Basic net income (loss) per share is calculated by dividing net income (loss) by the weighted-average number of common shares outstanding during the year. Diluted net income per share is calculated by using the weighted-average number of common shares outstanding during the year increased to include the number of additional shares of common stock that would have been outstanding if the dilutive potential shares of common stock had been issued. The dilutive effect of outstanding options, Employee Stock Purchase Plan shares and restricted stock units is reflected in diluted net income per share by application of the treasury stock method, which includes consideration of stock-based compensation.

For years presented with a net diluted net loss per common share is the same as basic net loss per common share, as the effect of the potential common stock equivalents is anti-dilutive and as such is excluded from the calculations of the diluted net loss per share.

The following table sets forth the computation of basic and diluted net loss and the weighted average number of shares used in computing basic and diluted net loss per share (in thousands):

	Year Ended December 31,						
	2010	2009	2008				
Numerator:							
Net loss—basic and diluted	\$ (10,518)	\$ (17,679)	\$ (2,869)				
Denominator:	,						
Weighted-average number of common shares							
outstanding used in computing basic net loss per							
share	13,540	13.279	12,770				
Dilutive potential common shares used in computing	,	10,000	12,770				
diluted net loss per share	_	·					
Total weighted-average number of shares used in	<u> </u>						
computing diluted net loss per share	13,540	13,279	12,770				
							

Anti-dilutive Securities

The following number of weighted shares outstanding, prior to the application of the treasury stock method, were excluded from the computation of diluted net loss per common share for the years presented because including them would have had an anti-dilutive effect (in thousands):

	Yea	r Ended December 31,	:	ert in
	2010	2009	200)8
Options to purchase common stock.	3,187	2,746		2,882
Restricted stock units	48	- 5		- 19
Employee stock purchase plan shares	66	84		94
Total	3,301	2,835		2,995

NOTE 9—DEFINED CONTRIBUTION PLAN:

In the United States, the Company has an employee savings plan (401(k) Plan) that qualifies as a deferred salary arrangement under Section 401(k) of the Internal Revenue Code. Eligible employees may make voluntary contributions to the 401(k) Plan up to 100% of their annual compensation, subject to statutory annual limitations. From April 1999 to December 31, 2008, the Company made discretionary matching contributions of 50% to 75% of all U.S. employees' contributions in each 401(k) Plan year. The Company made no discretionary contributions in 2010 and 2009 and made discretionary contributions of \$572,000 in 2008 under the 401(k) Plan.

For the Company's Japanese subsidiary, it has established an employee retirement plan at its discretion. In addition, for some of the Company's other foreign subsidiaries, the Company deposits funds with insurance companies, third-party trustees, or into government-managed accounts consistent with the requirements of local laws. The Company has fully funded or accrued for its obligations as of December 31, 2010, and the related expense for each of the three years then ended was not significant.

NOTE 10—SEGMENT, GEOGRAPHIC AND MAJOR CUSTOMER INFORMATION:

In accordance with the FASB ASC 280 guidance on disclosures about segments of an enterprise and related information, operating segments are identified as components of an enterprise about which separate discrete financial information is available for evaluation by the chief operating decision maker, or decision-making group, in making decisions how to allocate resources and assess performance. The Company's chief decision maker, as defined under the FASB's ASC 280 guidance, is a combination of the Chief Executive Officer and the Executive Vice President and Chief Financial Officer. To date, the Company has viewed its operations, managed its business, and used one measurement of profitability for the one operating segment – the sale of aesthetic medical equipment and services, and distribution of cosmeceutical and dermal filler products, to qualified medical practitioners. In addition, substantially all of the Company's long-lived assets are located in one facility in the United States. As a result, the financial information disclosed in the Company's Consolidated Financial Statements represents all of the material financial information related to the Company's operating segment. The following table summarizes revenue by geographic region, which is based on the shipping location of where the product is delivered, and product category (in thousands):

	Year Ended December 31,			
	2010	2009(1)	2008(1)	
Revenue mix by geography:		. —		
United States	\$ 19,337	\$ 21,019	\$ 41,683	
Japan	13,625	9,636	10,929	
Asia, excluding Japan	5,131	4,727	5,713	
Europe	5,801	7,087	10,522	
Rest of the world	9,380	11,213	14,532	
Consolidated total	\$ 53,274	\$ 53,682	\$ 83,379	
Revenue mix by product category:				
Products	\$ 27,808	\$ 26,842	\$ 57,821	
Upgrades	4,824	6,343	8,361	
Service	13,231	13,186	11,358	
Titan hand piece refills	3,863	5,599	5,662	
Dermal filler and cosmeceuticals ⁽¹⁾	3,548	1,712	177	
Consolidated total	\$ 53,274	\$ 53,682	\$ 83,379	

⁽¹⁾ Beginning in 2010, we classified revenue from Dermal fillers and cosmeceuticals product in the revenue category 'Dermal fillers and cosmeceuticals.' Previously, we classified this revenue under the category of 'Products.' As such, we reclassified the 2009 and 2008 revenue from 'Products' to 'Dermal fillers and cosmeceuticals.'

The Company had one customer that represented net revenue of 5% in 2010, 7% in 2009 and 14% in 2008.

The Company had one customer that accounted for 10% at December 31, 2010 and 29% at December 31, 2009 of the Company's total accounts receivable balance.

NOTE 11—COMMITMENTS AND CONTINGENCIES:

Facility Leases.

The Company leases its Brisbane, California, office and manufacturing facility under a non-cancelable operating lease which expires in 2017. In addition, the Company has leased office facilities in certain international countries as follows:

Country	Square Footage	Lease termination or Expiration
Japan	Approximately 5,790	Three leases, of which two expire in May 2012, and one expires in July 2013, but may be cancelled at any time with a six-month notice.
Switzerland	Approximately 3,174	One lease expiring March 2013.
France	Approximately 450	Lease expires in November 2011, but may be cancelled at any time with a three-month notice.
Śpain	Approximately 175	Lease automatically renews at the end of each six-month period.

As of December 31, 2010, the Company was committed to minimum lease payments for facilities and other leased assets under long-term non-cancelable operating leases as follows (in thousands):

Year Ending December 31,	 Amount
2011	\$ 1,688
2012	1,509
2013	1,281
2014	1,232
2015	1,269
2016 and thereafter	2,653
Total Future minimum rental payments	\$ 9,632

Gross rent expense was \$1.7 million in 2010, \$1.6 million in 2009 and \$1.7 million in 2008.

Purchase Commitments.

The Company maintains certain open inventory purchase commitments with its suppliers to ensure a smooth and continuous supply for key components. The Company's liability in these purchase commitments is generally restricted to a forecasted time-horizon as agreed between the parties. These forecasted time-horizons can vary among different suppliers. The Company's open inventory purchase commitments with its suppliers were not significant at December 31, 2010.

Indemnifications

In the normal course of business, the Company enters into agreements that contain a variety of representations, warranties, and indemnification obligations. For example, the Company has entered into indemnification agreements with each of its directors and executive officers. The Company's exposure under its various indemnification obligations is unknown and not reasonably estimable as they involve future claims that may be made against the Company. As such, the Company has not accrued any amounts for such obligations.

Litigation

A Telephone Consumer Protection Act, or TCPA, class action lawsuit was filed against the Company in January 2008 and settled in 2009 on a class-wide basis. In 2009, the Company paid a total of \$950,000 in exchange for a full release of all claims and recorded a charge of \$850,000 in its 2009 Consolidated Statements of Operations for the cost of the settlement, net of the administrative expenses and contributions from its insurance carrier.

Other Legal Matters

In addition to the foregoing lawsuits, the Company is named from time to time as a party to product liability and contractual lawsuits in the normal course of its business. As of December 31, 2010, the Company was not a party to any material pending litigation other than those described above in the "Litigation" section.

NOTE 12—SUBSEQUENT EVENT:

Management evaluated all activity of the Company and concluded that no subsequent events have occurred that would require recognition in the Consolidated Financial Statements or disclosure in Notes to Consolidated Financial Statements as of December 31, 2010.

SUPPLEMENTARY FINANCIAL DATA (UNAUDITED) (In thousands, except per share amounts)

Quarter ended:	Dec. 31, 2010	Sept. 30, 2010	June 30, 2010	March 31, 2010	Dec. 31, 2009	Sept. 30, 2009	June 30, 2009	March 31, 2009
2 .	\$15,216	\$12,092	\$12,217	\$ 13,749	\$15,416	\$ 12,171	\$11,665	\$ 14,430
Cost of revenue	6,233	5,661	5,335	5,829	5,783	4,910	5,130	5,936
Gross profit	8,983	6,431	6,882	7,920	9,633	7,261	6,535	8,494
Operating expenses:								
Sales and marketing	6,123	5,799	6,452	6,361	6,100	5,112	6,071	7,003
Research and development	2,173	1,871	1,506	1,454	1,888	1,684	1,495	1,743
General and administrative	2,238	2,352	2,744	2,242	2,063	2,121	3,616	2,520
Litigation settlement								850
Total operating expense	10,534	10,022	10,702	10,057	10,051	8,917	11,182	12,116
Loss from operations	(1,551)	(3,591)	(3,820)	(2,137)	(418)	(1,656)	(4,647)	(3,622)
Interest and other income, net.	144	132	<u> </u>	<u>166</u>	174	288	511	599
Loss before income taxes	(1,407)	(3,459)	(3,679)	(1,971)	(244)	(1,368)	(4,136)	(3,023)
Provision (benefit) for income								
taxes	(127)		82	47	(251)	12,126	(1,772)	(1,195)
Net income (loss)	\$ (1,280)	\$ (3,459)	\$ (3,761)	\$ (2,018)	<u>\$ 7</u>	\$ (13,494)	\$ (2,364)	<u>\$ (1,828)</u>
Net loss per share—basic	\$ (0.09)	\$ (0.25)	\$ (0.28)	\$ (0.15)	\$ 0.00	<u>\$ (1.01)</u>	\$ (0.18)	\$ (0.14)
Net loss per share—diluted	\$ (0.09)	\$ (0.25)	\$ (0.28)	\$ (0.15)	\$ 0.00	\$ (1.01)	\$ (0.18)	\$ (0.14)
Weighted average number of shares used in per share calculations:				: •				
Basic	13,622	13,589	13;501	13,438	13,427	13,382	13,317	13,120
Diluted	13,622	13,589	13,501	13,438	13,610	13,382	13,317	13,120

SCHEDULE II

CUTERA, INC.

VALUATION AND QUALIFYING ACCOUNTS (in thousands) For the Year Ended December 31, 2010, 2009 and 2008

	В	Salance at Seginning of Year	A	dditions	Dedu	ıctions		Balance at End of Year
Allowance for doubtful accounts receivable						;		4 1 1
Year ended December 31, 2010	\$	586	\$	116	\$	682	\$	20
Year ended December 31, 2009	\$	61	\$	675	\$	150	\$	586
Year ended December 31, 2008	\$	9	\$	191	\$	139	: .\$	61

ITEM 9. CHANGES IN AND DISAGREEMENTS WITH ACCOUNTANTS ON ACCOUNTING AND FINANCIAL DISCLOSURE

None.

ITEM 9A. CONTROLS AND PROCEDURES

Evaluation of Disclosure Controls and Procedures

Attached as exhibits to this Annual Report are certifications of the Company's Chief Executive Officer (CEO) and Chief Financial Officer (CFO), which are required in accordance with Rule 13a-14 of the Securities Exchange Act of 1934, as amended (Exchange Act). This Controls and Procedures section includes the information concerning the controls evaluation referred to in the certifications, and it should be read in conjunction with the certifications for a more complete understanding of the topics presented.

The Company conducted an evaluation of the effectiveness of the design and operation of its disclosure controls and procedures (as defined in the Rules 13a-15(e) and 15d-15(e) under the Exchange Act) (Disclosure Controls) as of the end of the period covered by this Report required by Exchange Act Rules 13a-15(b) or 15d-15(b). The controls evaluation was conducted under the supervision and with the participation of the Company's management, including the CEO and CFO. Based on this evaluation, the CEO and our CFO have concluded that as of the end of the period covered by this report the Company's disclosure controls and procedures were effective at a reasonable assurance level.

Definition of Disclosure Controls

Disclosure Controls are controls and procedures designed to reasonably assure that information required to be disclosed in the Company's reports filed under the Exchange Act, such as this Report, is recorded, processed, summarized and reported within the time periods specified in the SEC's rules and forms. Disclosure Controls are also designed to reasonably assure that such information is accumulated and communicated to the Company's management, including the CEO and CFO, as appropriate to allow timely decisions regarding required disclosure. The Company's Disclosure Controls include components of its internal control over financial reporting, which consists of control processes designed to provide reasonable assurance regarding the reliability of its financial reporting and the preparation of financial statements in accordance with generally accepted accounting principles in the U.S. To the extent that components of the Company's internal control over financial reporting are included within its Disclosure Controls, they are included in the scope of the Company's annual controls evaluation.

Management's Report on Internal Control over Financial Reporting

The Company's management is responsible for establishing and maintaining adequate internal control over financial reporting as defined in Rules 13a-15(f) and 15d-15(f) under the Exchange Act. Under the supervision and with the participation of the Company's management, including the CEO and CFO, the Company conducted an evaluation of the effectiveness of its internal control over financial reporting based on criteria established in the framework in Internal Control—Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission. Based on this evaluation, the Company's management concluded that the Company's internal control over financial reporting was effective as of December 31, 2010. The effectiveness of our internal control over financial reporting as of December 31, 2010 has been audited by PricewaterhouseCoopers LLP, an Independent Registered Public Accounting Firm, as stated in their report, which is included herein.

Limitations on the Effectiveness of Controls

The Company's management, including the CEO and CFO, does not expect that the Company's disclosure controls or internal control over financial reporting will prevent all error and all fraud. A control system, no matter how well designed and operated, can provide only reasonable, not absolute, assurance that the control system's objectives will be met. Further, the design of a control system must reflect the fact that there are resource constraints, and the benefits of controls must be considered relative to their costs. Because of the inherent limitations in all control systems, no evaluation of controls can provide absolute assurance that all control issues and instances of fraud, if any, within the Company have been detected. These inherent limitations include the realities that judgments in decision making can be faulty and that breakdowns can occur because of simple error or mistake. Controls can also be circumvented by the individual acts of some persons, by collusion of two or more people, or by management override of the controls. The design of any system of controls is based in part on certain assumptions about the likelihood of future events, and there can be no assurance that any design will succeed in achieving its stated goals under all potential future conditions. Over time, controls may become inadequate because of changes in conditions or deterioration in the degree of compliance with policies or procedures. Because of the inherent limitations in a cost-effective control system, misstatements due to error or fraud may occur and not be detected.

Changes in Internal Control over Financial Reporting

There were no changes in the Company's internal control over financial reporting that occurred during the most recent fiscal quarter that have materially affected, or are reasonably likely to materially affect, the Company's internal control over financial reporting.

ITEM 9B. OTHER INFORMATION

The Company has established that the 2011 Annual Meeting of Stockholders will be held at its principal executive offices located at 3240 Bayshore Blvd., Brisbane, CA 94005-1021 on June 13, 2011 at 10:00 a.m. and the record date for the purposes of voting in that meeting shall be April 18, 2011.

PART III

Certain information required by Part III is omitted from this Annual Report on Form 10-K because we will file a Definitive Proxy Statement (the "Proxy Statement") for our 2011 Annual Meeting of Stockholders with the Securities and Exchange Commission within 120 days after the end of our fiscal year ended December 31, 2010.

ITEM 10. DIRECTORS, EXECUTIVE OFFICERS AND CORPORATE GOVERNANCE

The information required by this Item is incorporated herein by reference to the Proxy Statement.

ITEM 11. EXECUTIVE COMPENSATION

The information required by this Item is incorporated herein by reference to the Proxy Statement.

ITEM 12. SECURITY OWNERSHIP OF CERTAIN BENEFICIAL OWNERS AND MANAGEMENT AND RELATED STOCKHOLDER MATTERS

The information required by this Item is incorporated herein by reference to the Proxy Statement.

ITEM 13. CERTAIN RELATIONSHIPS AND RELATED TRANSACTIONS, AND DIRECTOR INDEPENDENCE

The information required by this Item is incorporated herein by reference to the Proxy Statement.

ITEM 14. PRINCIPAL ACCOUNTING FEES AND SERVICES

The information required by this Item is incorporated herein by reference to the Proxy Statement.

PART IV

ITEM 15. EXHIBITS AND FINANCIAL STATEMENT SCHEDULES

- (1) The financial statements required by Item 15(a) are filed as Item 8 of this annual report.
- (2) The financial statement schedule required by Item 15(a) filed as Item 8 of this annual report.
- (3) Exhibits.

Exhibit No.	Description
3.2 ⁽¹⁾	Amended and Restated Certificate of Incorporation of the Registrant (Delaware).
3.4(1)	Bylaws of the Registrant.
4.1 ⁽⁴⁾	Specimen Common Stock certificate of the Registrant.
10.1(1)	Form of Indemnification Agreement for directors and executive officers.
$10.2^{(1)}$	1998 Stock Plan.
10.3(1)	2004 Equity Incentive Plan.
$10.4^{(5)}$	2004 Employee Stock Purchase Plan.
$10.6^{(1)}$	Brisbane Technology Park Lease dated August 5, 2003 by and between the Registrant and Gal-
	Brisbane, L.P. for office space located at 3240 Bayshore Boulevard, Brisbane, California.
$10.10^{(2)}$	Settlement Agreement and Non-Exclusive Patent License, each between the Registrant and Palomar
	Medical Technologies, Inc. dated June 2, 2006.
$10.11^{(3)}$	Form of Performance Unit Award Agreement.
10.13 ^{(4)†}	Distribution Agreement between the Registrant and PSS World Medical Shared Services, Inc., a
	subsidiary of PSS World Medical dated October 1, 2006.
10.14 ⁽⁶⁾	Cutera, Inc. 2004 Equity Incentive Plan, as amended by its Board of Directors on April 25, 2008.
$10.18^{(7)}$	Consulting Agreement dated March 2, 2009 by and between the Company and David A. Gollnick.
$10.19^{(8)}$	First Amendment to Brisbane Technology Park Lease dated August 11, 2010 by and between the
	Company and BMR-Bayshore Boulevard LLC, as successor-in-interest to Gal-Brisbane, L.P., the
	original landlord, for office space located at 3240 Bayshore Boulevard.
10.20	Change of Control and Severance Agreement dated January 5, 2011 by and between the Company and
	Len DeBenedictis, Chief Technology Officer of Cutera Inc.
23.1	Consent of Independent Registered Public Accounting Firm.
24.1	Power of Attorney (see page 73).
31.1	Certification of Chief Executive Officer pursuant to Section 302 of the Sarbanes-Oxley Act of 2002.
31.2	Certification of Chief Financial Officer pursuant to Section 302 of the Sarbanes-Oxley Act of 2002.
32.1	Certification of Chief Executive Officer and Chief Financial Officer pursuant to 18 U.S.C. Section
	1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002.

- (1) Incorporated by reference from our Registration Statement on Form S-1 (Registration No. 333-111928) which was declared effective on March 30, 2004.
- (2) Incorporated by reference from our Current Report on Form 8-K filed on June 2, 2006.
- (3) Incorporated by reference from our Quarterly Report on Form 10-Q filed on November 14, 2005.
- (4) Incorporated by reference from our Quarterly Report on Form 10-Q filed on November 8, 2006.
- (5) Incorporated by reference from our 2006 Annual Report on Form 10-K filed on March 16, 2007.
- (6) Incorporated by reference from our Definitive Proxy Statement on Form 14A filed with the SEC on April 28, 2008.
- (7) Incorporated by reference from our Current Report on Form 8-K filed on March 4, 2009.
- (8) Incorporated by reference from our Quarterly Report on Form 10-Q filed on November 1, 2010.
- † Confidential Treatment has been requested for certain portions of this exhibit.

SIGNATURES

Pursuant to the requirements of Section 13 or 15(d) of The Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized, in the city of Brisbane, State of California, on the 15th day of March, 2011.

CUTERA, INC.

By:	/s/ KEVIN P. CONNORS
	Kevin P. Connors
	President and Chief Executive Officer

Power of Attorney

KNOW ALL MEN AND WOMEN BY THESE PRESENTS, that each person whose signature appears below constitutes and appoints Kevin P. Connors, his attorney-in-fact, for him or her in any and all capacities, to sign any amendments to this Annual Report on Form 10-K, and to file the same, with exhibits thereto and other documents in connection therewith, with the U.S. Securities and Exchange Commission, hereby ratifying and confirming all that said attorney-in-fact, or his substitute, may do or cause to be done by virtue thereof.

Pursuant to the requirements of the Securities Exchange Act of 1934, this report has been signed below by the following persons on behalf of the registrant and in the capacities and on the dates indicated.

Signature	Title	Date
/s/ KEVIN P. CONNORS Kevin P. Connors	President, Chief Executive Officer and Director (Principal Executive Officer)	March 15, 2011
/s/ RONALD J. SANTILLI Ronald J. Santilli	Executive Vice President and Chief Financial Officer (Principal Accounting Officer)	March 15, 2011
/s/ DAVID B. APFELBERG David B. Apfelberg	Director	March 15, 2011
/s/ DAVID A. GOLLNICK David A. Gollnick	Director	March 15, 2011
/s/ MARK LORTZ Mark Lortz	Director	March 15, 2011
/s/ TIM O'SHEA Tim O'Shea	Director	March 15, 2011
/s/ JERRY P. WIDMAN Jerry P. Widman	Director	March 15, 2011

CERTIFICATION OF CHIEF EXECUTIVE OFFICER PURSUANT TO 15 U.S.C. SECTION 7241, AS ADOPTED PURSUANT TO SECTION 302 OF THE SARBANES-OXLEY ACT OF 2002

I, Kevin P. Connors, certify that:

- 1. I have reviewed this annual report on Form 10-K of Cutera, Inc.:
- 2. Based on my knowledge, this report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this report;
- 3. Based on my knowledge, the financial statements, and other financial information included in this report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this report;
- 4. The registrant's other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - (a) Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this annual report is being prepared;
 - (b) Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - (c) Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this report based on such evaluation; and
 - (d) Disclosed in this report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
- 5. The registrant's other certifying officer and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors (or persons performing the equivalent functions):
 - a) All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - b) Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

(Principal Executive Officer)

Date: March 15, 2011

/s/ KEVIN P. CONNORS

Kevin P. Connors

President, Chief Executive Officer and Director

CERTIFICATION OF CHIEF FINANCIAL OFFICER PURSUANT TO 15 U.S.C. SECTION 7241, AS ADOPTED PURSUANT TO SECTION 302 OF THE SARBANES-OXLEY ACT OF 2002

I, Ronald J. Santilli, certify that:

- 1. I have reviewed this annual report on Form 10-K of Cutera, Inc.:
- 2. Based on my knowledge, this report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this report;
- 3. Based on my knowledge, the financial statements, and other financial information included in this report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this report;
- 4. The registrant's other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - (a) Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this annual report is being prepared;
 - (b) Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - (c) Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this report based on such evaluation; and
 - (d) Disclosed in this report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
- 5. The registrant's other certifying officer and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors or (any other persons performing the equivalent functions):
 - a) All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - b) Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

March 15, 2011

/s/ RONALD J. SANTILLI

Ronald J. Santilli

Chief Financial Officer and Executive Vice President
(Principal Financial and Accounting Officer)

CERTIFICATION OF CHIEF EXECUTIVE OFFICER AND CHIEF FINANCIAL OFFICER PURSUANT TO 18 U.S.C. SECTION 1350, AS ADOPTED PURSUANT TO SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002

In connection with the annual report on Form 10-K of Cutera, Inc. a Delaware corporation, for the period ended December 31, 2010, as filed with the Securities and Exchange Commission, each of the undersigned officers of Cutera, Inc. certifies pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002, that to his respective knowledge:

- (1) the annual report fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934, as amended; and
- the information contained in the annual report fairly presents, in all material respects, the financial condition and results of operations of Cutera, Inc. for the periods presented therein.

Date: March 15, 2011 /s/ Kevin P. Connors

Kevin P. Connors
President, Chief Executive Officer and Director
(Principal Executive Officer)

Date: March 15, 2011 /s/ Ronald J. Santilli

Ronald J. Santilli Chief Financial Officer and Executive Vice President (Principal Financial and Accounting Officer)

Corporate Information (as of April 26, 2011)

BOARD OF DIRECTORS

Kevin P. Connors, President and Chief Executive Officer, Cutera, Inc.

David A. Gollnick^I, Former Vice President of Research and Development, Cutera, Inc.

David B. Apfelberg, MD^{3 5}, Clinical Professor of Plastic Surgery, Stanford University Medical Center

Mark Lortz², Former Chief Executive Officer, ThereaSense, Inc.

Timothy J. O'Shea^{2 3}, Managing Director, Oxo Capital

Jerry P. Widman ²³⁴, Former Chief Financial Officer, Ascension Health 1-Mr. Gollnick is a member of our Board of Directors and provides consulting services to the Company.

- 2-Audit Committee member
- 3-Compensation Committee member
- 4-Chairman of Audit Committee
- 5-Chairman of Compensation Committee

MANAGEMENT TEAM

Kevin P. Connors, President, Chief Executive Officer and Director

Len DeBenedictis, Chief Technology Officer

Ronald J. Santilli, Executive Vice President and Chief Financial Officer Scott Davenport, Vice President.

Research and Development

Brian Hall, Vice President, Service

Michael Poole, Vice President, North

America Sales

Robert Shine, Vice President, Technical

Marketing

Chris West, Vice President, Pacific Rim Ken Witte, Vice President, Marketing & Clinical Research

ANNUAL MEETING

Annual meeting of stockholders will be held on June 14, 2011, 10:00 a.m. (PDT) at: 3240 Bayshore Blvd., Brisbane, California 94005.

TRANSFER AGENT Computershare Trust Company, Inc. 350 Indiana St., Suite 800 Golden, Colorado 80401 303-262-0600

INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM PricewaterhouseCoopers LLP San Jose, California

CORPORATE LEGAL COUNSEL Wilson, Sonsini, Goodrich & Rosati, P.C., Palo Alto, California

CORPORATE/STOCKHOLDER INFORMATION

Our Form 10-K was filed with the Securities and Exchange Commission on March 15, 2011. For additional copies of this report, Form 10-K, or other financial information, without charge, please visit the Investor Relations page on our website at: www.cutera.com or write to ir@cutera.com.

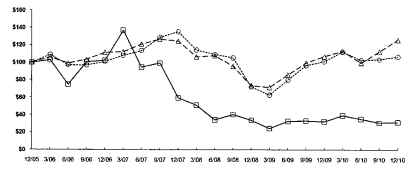
STOCK LISTING AND MARKET DATA

Our common stock is traded on The NASDAQ Global market under the symbol "CUTR." We have not declared or paid any cash dividends on our capital stock since our inception. We currently expect to retain future earnings, if any, for use in the operation and expansion of our business and do not anticipate paying any cash dividends in the foreseeable future. As of February 28, 2011, we believe there were approximately 2,600 holders of record of our common stock. The following table sets forth quarterly high and low closing sales prices per share of our common stock as reported on The NASDAQ Global Market for the periods indicated.

		•	Co	mmo	n S	Stock				
	Ξ	2010				2009				
		High		Low		High	_]	Low		
4th Qtr	\$	8.39	\$	7.01	\$	9.63	\$	7.97		
3rd Qtr		9.00		6.99		9.40		7.85		
2nd Qtr		12.04		8.62		9.03		5.93		
1st Qtr		11.03		8.25		8.71		5.57		

COMPARISON OF 5 YEAR CUMULATIVE TOTAL RETURN*

Among Cutera, Inc., the NASDAQ Composite Index and the NASDAQ Medical Equipment Index



*\$100 invested on 12/31/05 in stock or index, including reinvestment of dividends. Fiscal year ending December