

# INSPIREMD, INC.

## FORM 8-K (Current report filing)

Filed 05/12/15 for the Period Ending 05/12/15

Address	321 COLUMBUS AVENUE BOSTON, MA 02116
Telephone	(857) 453-6553
CIK	0001433607
Symbol	NSPR
SIC Code	3841 - Surgical and Medical Instruments and Apparatus
Industry	Medical Equipment & Supplies
Sector	Healthcare
Fiscal Year	12/31

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

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**FORM 8-K**

**CURRENT REPORT**  
**Pursuant to Section 13 or 15(d) of the**  
**Securities Exchange Act of 1934**

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Date of Report (Date of earliest event reported): May 12, 2015

**InspireMD, Inc.**

(Exact name of registrant as specified in its charter)

Delaware  
(State or other  
jurisdiction  
of incorporation)

001-35731  
(Commission File Number)

26-2123838  
(IRS Employer  
Identification No.)

321 Columbus Avenue  
Boston, Massachusetts  
(Address of principal executive offices)

02116  
(Zip Code)

Registrant's telephone number, including area code: (857) 453-6553

(Former name or former address, if changed since last report)

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Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
  - Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
  - Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
  - Pre-commencement communications pursuant to Rule 13e-4 (c) under the Exchange Act (17 CFR 240.13e-4(c))
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**Item 8.01 Other Events.**

On May 12, 2015, InspireMD, Inc. issued a press release announcing that there will be multiple clinical presentations across a broad range of indications for MicroNet™ Technology at the EuroPCR 2015 Conference to be held from May 19 to May 22, 2015, in Paris, France.

A copy of the press release is attached as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated by reference herein.

**Item 9.01 Financial Statements and Exhibits.**

(d) Exhibits

<b>Exhibit Number</b>	<b>Description</b>
99.1	Press release dated May 12, 2015

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**SIGNATURES**

Pursuant to the requirements of the Securities Exchange Act of 1934, as amended, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

**INSPIREMD, INC.**

Date: May 12, 2015

By: /s/ Craig Shore

Name: Craig Shore

Title: Chief Financial Officer

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## InspireMD Announces Clinical Presentations on CGuard and MGuard Systems with MicroNet™ Technology at EuroPCR 2015 Conference

*Clinical Presentations Highlight Carotid and Coronary Applications of InspireMD's Proprietary MicroNet™ Technology*

**BOSTON, MA – May 12, 2015** – InspireMD, Inc. (NYSE MKT: NSPR) (“InspireMD” or the “Company”), a leader in stent embolic protection systems (“EPS”), today announced that there will be multiple clinical presentations across a broad range of indications for MicroNet™ Technology at the upcoming EuroPCR conference from May 19-22, 2015 in Paris, France.

Each year, more than 120 companies from the cardiovascular industry, including device and equipment manufacturers, attend EuroPCR, the leading cardiovascular event in Europe. This event allows attendees to discover new products and R&D projects, as well as interact with practitioners and industry partners to drive continued development and innovation in the cardiovascular field.

Alan Milinazzo, CEO of InspireMD, commented, “EuroPCR is a major annual meeting for InspireMD and we have a very busy schedule of meetings with customers, distributors and potential future partners ahead. We look forward to productive sessions and making progress on our many initiatives.”

Milinazzo continued, “Feedback on our CGuard Embolic Prevention system has been positive, with solid physician interest and the device steadily gaining market traction. We look forward to the clinical presentations at EuroPCR 2015 from a range of physicians, including Dr. Piotr Musialek of Poland and Dr. Joachim Schofer of Germany.”

At EuroPCR 2015, there will be 7 clinical presentations featuring InspireMD’s MicroNet™ Technology, with applications in the carotid, coronary arteries, and saphenous vein grafts. The sessions include:

### **Wednesday, May 20<sup>th</sup>**

09:20 – 10:20, Room 243

#### ***Carotid and Supra-Aortic Disease Management***

Mesh-covered stent in routine endovascular management of thrombus containing/subtotal lesions: a new paradigm in carotid revascularization?  
*Piotr Musialek, Poland*

15:45 – 16:45, Room Ternes 2

#### ***Interactive Case Corner: Challenging Peripheral Vascular Interventions***

Mesh-covered stent in endovascular management of highly calcific lesions: a new paradigm in carotid revascularization? *Adam Mazurek, Poland*

### **Thursday, May 21<sup>st</sup>**

08:15 – 09:15, Room 241

#### ***Primary PCI: Technical and Strategic Challenges***

In-hospital and long-term outcomes of mesh-covered MGuard stent implantation for treatment of STEMI with high thrombus burden despite manual aspiration. *Enrico Cerrato, Italy*

12:30 – 13:30, Room 343

#### ***Solutions to Treat Degenerative Saphenous Vein Grafts***

When a mesh-covered stent saves the day. *Alexander Goldberg, Israel*



**Friday, May 22<sup>nd</sup>**

10:40 – 11:40, Room 353

***Coronary Aneurysms after PCI***

STEMI patient and stent fracture treated with covered stent. *David Baghdasaryan, Armenia*

11:45 – 11:52, Room 242A

***Evolutions in Carotid Angioplasty***

Prospective, multicentre study on the safety and efficacy of a novel mesh-covered carotid stent in patients with symptomatic and asymptomatic carotid artery stenosis: the CGuard CARotid Embolic protection using microNET trial (CARENET). *Joachim Schofer, Germany*

12:09 – 12:16, Room 242A

***Evolutions in Carotid Angioplasty***

Novel PARADIGM in carotid revascularization: prospective evaluation of all-comer percutaneous carotid revascularization in symptomatic and increased-risk asymptomatic carotid artery stenosis using CGuard mesh-covered embolic prevention stent system. *Piotr Musialek, Poland*

For more information about InspireMD and its offerings, visit [www.inspiremd.com](http://www.inspiremd.com).

**About InspireMD, Inc.**

InspireMD seeks to utilize its proprietary MGuard™ with MicroNet™ technology to make its products the industry standard for embolic protection and to provide a superior solution to the key clinical issues of current stenting in patients with a high risk of distal embolization, no reflow and major adverse cardiac events.

InspireMD intends to pursue applications of this MicroNet technology in coronary, carotid (CGuard™), neurovascular, and peripheral artery procedures. InspireMD's common stock is quoted on the NYSE MKT under the ticker symbol NSPR.

**Forward-looking Statements:**


*This press release contains "forward-looking statements." Such statements may be preceded by the words "intends," "may," "will," "plans," "expects," "anticipates," "projects," "predicts," "estimates," "aims," "believes," "hopes," "potential" or similar words. Forward-looking statements are not guarantees of future performance, are based on certain assumptions and are subject to various known and unknown risks and uncertainties, many of which are beyond the Company's control, and cannot be predicted or quantified and consequently, actual results may differ materially from those expressed or implied by such forward-looking statements. Such risks and uncertainties include, without limitation, risks and uncertainties associated with (i) market acceptance of our existing and new products, (ii) negative clinical trial results or lengthy product delays in key markets, (iii) an inability to secure regulatory approvals for the sale of our products, (iv) intense competition in the medical device industry from much larger, multinational companies, (v) product liability claims, (vi) product malfunctions, (vii) our limited manufacturing capabilities and reliance on subcontractors for assistance, (viii) insufficient or inadequate reimbursement by governmental and other third party payers for our products, (ix) our efforts to successfully obtain and maintain intellectual property protection covering our products, which may not be successful, (x) legislative or regulatory reform of the healthcare system in both the U.S. and foreign jurisdictions, (xi) our reliance on single suppliers for certain product components, (xii) the fact that we will need to raise additional capital to meet our business requirements in the future and that such capital raising may be costly, dilutive or difficult to obtain and (xiii) the fact that we conduct business in multiple foreign jurisdictions, exposing us to foreign currency exchange rate fluctuations, logistical and communications challenges, burdens and costs of compliance with foreign laws and political and economic instability in each jurisdiction. More detailed information about the Company and the risk factors that may affect the realization of forward looking statements is set forth in the Company's filings with the Securities and Exchange Commission (SEC), including the Company's Annual Report on Form 10-K and its Quarterly Reports on Form 10-Q. Investors and security holders are urged to read these documents free of charge on the SEC's web site at <http://www.sec.gov>. The Company assumes no obligation to publicly update or revise its forward-looking statements as a result of new information, future events or otherwise.*



**Investor Contacts:**

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