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

FORM 8-K

Current Report Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): April 1, 2019

DARÉ BIOSCIENCE, INC. (Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation)

 $Pre-commencement communications \ pursuant \ to \ Rule \ 13e-4(c) \ under \ the \ Exchange \ Act \ (17 \ CFR \ 240.13e-4(c))$

001-36395

20-4139823

(Commission File Number)

(I.R.S. Employer Identification No.)

3655 Nobel Drive, Suite 260 San Diego, CA 92122 (Address of Principal Executive Offices and Zip Code)

Registrant's telephone number, including area code: (858) 926-7655

Not Applicable

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

hec	k 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 (see General Instruction A.2. below):
	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)
٦.	Pre-commencement communications pursuant to Rule 144-2(h) under the Eychange Act (17 CER 240 144-2(h))

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. x

Item 8.01 Other Events

Compliance with Nasdaq Minimum Bid Price Rule

On April 1, 2019, Daré Bioscience, Inc. ("Daré") was informed by the Listing Qualifications Department (the "Staff") of The Nasdaq Stock Market LLC that Daré regained compliance with the minimum bid price requirement in Nasdaq Listing Rule 5550(a)(2) as a result of the closing bid price of Daré's common stock being \$1.00 per share or greater for the 10 consecutive business day period from March 18, 2019 to March 29, 2019. As previously reported, Daré was not in compliance with the minimum bid price requirement because the closing bid price for its common stock was less than \$1.00 for the 30 consecutive business days preceding November 30, 2018. That matter is now closed.

Daré issued a press release regarding it regaining compliance with the minimum bid price requirement, a copy of which is included as Exhibit 99.1 to this report.

Corporate Presentation

Included as Exhibit 99.2 to this report is a presentation about Daré and its product candidates, dated April 1, 2019, which is incorporated herein by reference. Daré intends to use the presentation and its contents in various meetings with investors, securities analysts and others, commencing on April 1, 2019.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

99.2

Description

Press release dated April 1, 2019
Corporate presentation, dated April 1, 2019 99.1

SIGNATURES

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

DARÉ BIOSCIENCE, INC.

Dated: April 1, 2019 By: /s/ Sabrina Martucci Johnson

Name: Sabrina Martucci Johnson

Title: President and Chief Executive Officer

Daré Bioscience, Inc. Regains Listing Compliance on Nasdaq

SAN DIEGO, April 1, 2019 (GLOBE NEWSWIRE) -- Daré Bioscience, Inc.(NASDAQ: DARE), a leader in women's health innovation, today announced that it received notice from The Nasdaq Stock Market LLC indicating that the Company regained compliance with the minimum bid price requirement in Nasdaq Listing Rule 5550(a)(2) as a result of the closing bid price of the Company's common stock being \$1.00 per share or greater for the 10 consecutive business day period from March 18, 2019 to March 29, 2019 and that the previously reported non-compliance listing matter is now closed.

About Daré Bioscience

Daré Bioscience is a clinical-stage biopharmaceutical company committed to the advancement of innovative products for women's sexual health, vaginal health, fertility, and contraception. The company's mission is to identify, develop and bring to market a portfolio of novel, differentiated therapies that expand treatment options, improve outcomes and facilitate convenience for women in the areas of contraception, vaginal health, sexual health, and fertility.

Daré's product portfolio includes potential first-in-class candidates in clinical development: Ovaprene®, a non-hormonal, monthly contraceptive vaginal ring; Sildenafil Cream, 3.6%, a novel cream formulation of sildenafil to treat female sexual arousal disorder utilizing the active ingredient in Viagra®; DARE-BV1, a unique hydrogel formulation of clindamycin phosphate 2% to treat bacterial vaginosis via a single application; and DARE-HRT1, a combination bio-identical estradiol and progesterone intravaginal ring for hormone replacement therapy following menopause. To learn more about Daré's full portfolio of women's health product candidates, and mission to deliver novel therapies for women, please visit www.darebioscience.com.

Daré may announce material information about its finances, product candidates, clinical trials and other matters using its investor relations website (http://ir.darebioscience.com), SEC filings, press releases, public conference calls and webcasts. Daré will use these channels to distribute material information about the company, and may also use social media to communicate important information about the company, its finances, product candidates, clinical trials and other matters. The information Daré posts on its investor relations website or through social media channels may be deemed to be material information. Daré encourages investors, the media, and others interested in the company to review the information Daré posts on its investor relations website (https://darebioscience.gcs-web.com/) and to follow these Twitter accounts: @SabrinaDareCEO and @DareBioscience. Any updates to the list of social media channels the company may use to communicate information will be posted on the investor relations page of the company's website mentioned above.

Contacts:

Investors on behalf of Daré Bioscience, Inc.:

Alex Gray Burns McClellan agray@burnsmc.com 212-213-0006 OR

Media on behalf of Daré Bioscience, Inc.: Amanda Guisbond Canale Communications amanda@canalecomm.com 781-405-8775

Source: Daré Bioscience



Forward Looking Statements

THIS PRESENTATION IS FOR INFORMATIONAL PURPOSES ONLY AND IS NOT AN OFFER TO SELL OR A SOLICITATION OF AN OFFER TO BUY ANY SECURITIES OF DARÉ BIOSCIENCE, INC. ("DARÉ" OR THE "COMPANY"). THIS PRESENTATION INCLUDES CERTAIN INFORMATION OBTAINED FROM TRADE AND STATISTICAL SERVICES, THIRD PARTY PUBLICATIONS, AND OTHER SOURCES. DARÉ HAS NOT INDEPENDENTLY VERIFIED SUCH INFORMATION AND THERE CAN BE NO ASSURANCE AS TO ITS ACCURACY.

ALL STATEMENTS IN THIS PRESENTATION, OTHER THAN STATEMENTS OF HISTORICAL FACT, ARE FORWARD-LOOKING STATEMENTS WITHIN THE MEANING OF FEDERAL SECURITIES LAWS. IN SOME CASES, YOU CAN IDENTIFY FORWARD-LOOKING STATEMENTS BY TERM SUCH AS "MAY," "WILL," "EXPECT," "PLAN," "ANTICIPATE," "STRATEGY," "DESIGNED," "COULD," "INTEND," "BELIEVE," "ESTIMATE," "TARGET," ("POTENTIAL" AND OTHER SIMILAR EXPRESSIONS, OR THE NEGATIVE OF THESE TERMS. AS USED IN THIS PRESENTATION, "FIRST-IN-CATEGORY" IS A FORWARD-LOOKING STATEMENT REGARDING MARKET POTENTIAL OF A PRODUCT CANDIDATE. FORWARD-LOOKING STATEMENTS INVOLVE RISKS, UNCERTAINTIES AND ASSUMPTIONS THAT MAY CAUSE DARÉ'S ACTUAL RESULTS, PERFORMANCE OR ACHIEVEMENTS TO BE MATERIALLY DIFFERENT FROM THOSE EXPRESSED OR IMPLIED BY THE FORWARD-LOOKING STATEMENTS, INCLUDING, WITHOUT LIMITATION RISKS AND UNCERTAINTIES RELATING TO: THE OUTCOME OR SUCCESS OF CLINICAL TRIALS; DARÉ'S ABILITY TO RAISE ADDITIONAL CAPITAL AS NEEDED; DARÉ'S ABILITY TO OBTAIN AND MAINTAIN INTELLECTUAL PROPERTY PROTECTION FOR ITS PRODUCT CANDIDATES; DARÉ'S ABILITY TO DEVELOP PRODUCT CANDIDATES; DARÉ'S AND OTHER RISK FACTORS DESCRIBED IN DARÉ'S MOST RECENT ANNUAL REPORT ON FORM 10-K AND QUARTERLY REPORT ON FORM 10-Q FILED WITH THE SECURITIES AND EXCHANGE COMMISSION.

ALL FORWARD-LOOKING STATEMENTS IN THIS PRESENTATION ARE CURRENT ONLY AS OF THE DATE HEREOF AND DARÉ DOES NOT UNDERTAKE ANY OBLIGATION TO UPDATE ANY FORWARD-LOOKING STATEMENT TO REFLECT NEW INFORMATION, FUTURE DEVELOPMENTS OR OTHERWISE, EXCEPT AS REQUIRED BY LAW.



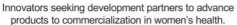
Vision: To become the premier innovation accelerator in women's health.

Mission: We achieve this by identifying, unlocking and advancing candidates with potential to be first-in-category, address persistent unmet needs and promote a better quality of life for women.



Accelerating Innovation in Women's Health

Market Misalignment is a Value Creation Opportunity



Large and mid-tier companies prefer to acquire or license produ that are later-stage or ready for commercialization.







Global Women's Health Market Worth \$51 Billion by 2025 - CAGR: 3.9% 1

Daring to be different

A pure play biopharmaceutical company focused on improving the health and well bei of women.

Our focus areas include:

- · Contraception / Pregnancy Prevention
- Sexual Health
- · Vaginal Health
- Fertility

Licensing, Partnering & Value Creation Strategy:

- · The portfolio is well positioned to drive upside
- · Product candidates are commercially viable and attractive to strategic partners
 - Products that have a data package including a proof-of-concept and/or the ability to leverage 505(b)(2) regulatory pathway
 - Candidates with the potential to be first-in-category that address persistent unmet needs i
 women's health
 - The ability to deliver products in a more *personalized* way for women



Daring to be different

We expect to deliver against multiple milestones over the next 12 - 24 months including:

- · Advancing our DARE-BV1 Bacterial Vaginosis (BV) program into a Phase 3 trial
- Topline readouts from our two pre-pivotal programs:
 - · Ovaprene (2H 2019) and
 - Sildenafil Cream, 3.6%, (4Q 2020)
- · Moving preclinical programs into Phase 1 development
 - DAER-HRT1 Hormone Replacement Therapy (HRT/VMS) Phase 1 (2019)
 - DARE-FRT1 Fertility and Pregnancy Maintenance (PTB/ART) Phase 1 (2020)
 - DARE-VVA1 –VVA in ER/PR+ Breast Cancer patients Phase 1 (2020)



Timeline reflects management's current estimates and constitutes a forward looking statement subject to qualifications elsewhere in the presentation. Actual development timeline may be substantially longer, and Daré is under n obligation to update or review this estimate.

Accelerating early-stage clinical programs with collaborations and non-dilutive funding whenever possible

Timeline reflects management's current estimates and constitutes a forward looking statement subject to qualifications elsewhere in the presentation. Actual development timeline may be substantially longer, and Daré is under robligation to update or review this estimate. "First-in-category" designations are forward looking statements based on currently available, FDA approved therapies.

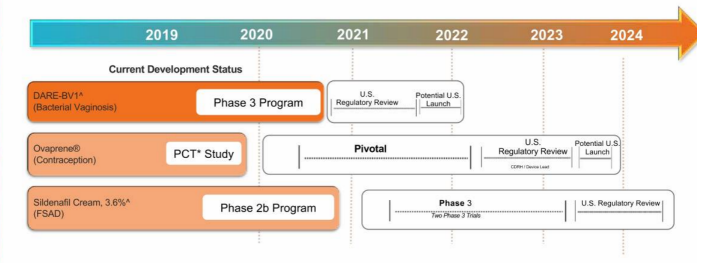
^505(b)(2) regulatory pathway anticipated.

*Ovaprene Post Coital Test (PCT) is a pre-pivotal clinical study.

*HRT Phase 1 study to be conducted in Australia by Daré subsidiary.



Portfolio Timeline Overview



Timeline reflects management's current estimates and constitutes a forward looking statement subject to qualifications elsewhere in the presentation. Actual development timeline may be substantially longer, and Daré is under no obligation to update or review this estimate.

'505(b)(2) regulatory pathway anticipated.
'Ovaprene Post Coital Test (PCT) is a pre-pivotal clinical study.





Program Overview



DARE-BV1 Overview

DARE-BV1

Phase 3 Initiation

Bacterial Vaginosis (BV)

Successful Proof of Concept

- Vaginal application of DARE-BV1 (clindamycin phosphate 2%) demonstrated effectiveness against BV in a proof-of-concept investigator initiated study in women (n=30):1
 - 88% of evaluable subjects met clinical cure endpoint at Test-of-Cure visit after single dose administered
 - Favorable efficacy profile over currently approved treatments

505(b)(2) Regulatory Pathway

· Single Phase 3 clinical trial planned for FDA approval

Attractive Market Opportunity

- BV is the most commonly reported vaginal infection in women ages 15-44 ²
- U.S. prevalence estimated to be ~21 million among women ages 14-49 ²
- Approved prescription drugs have less than optimal clinical cure rates (37-67%) 3
- Opportunity for significant upside and market expansion

Patent Coverage

- Patents covering the licensed technology have been granted with terms through 2028
- Additional patents pending would have terms through 2035

1. Data on file

2. https://www.cdc.gov/std/bv/stats.htm
3. BV Product Data: http://www.clindesse.com/pdf/Pl.pdf; http://www.accessdata.fda.gov/drugsatfda_docs/label/2014/205223s000lbl.pdf; http://www.accessdata.fda.gov/drugsatfda_docs/label/2014/205223s000lbl.pdf

Gel Delivery Technology

Features	Description	Innovative Product Profile			
In-Situ Gelation	Undergoes solution to gel (sol-to-gel) transition using body temperature as the trigger	 Allows product to be easily and directly placed at the site of infection Increased viscosity following application keeping the product at the site of application 			
Sustained-Erosion	Platform can be optimized to erode over a period of hours to multiple days	 Designed for a dual-release pattern providing maximal exposure time and amount of drug at the site of action Allows optimization of dosing duration for clindamycin – a time dependent antibiotic 			
Bio-Resorption and Adhesion	Hydrophilic ingredients are compatible with a variety of APIs	 Reinforces ability of product to bio-adhere at the site of application Eliminates need to remove product following completion of treatment regimen 			



Bacterial Vaginosis DARE-BV1 Proof of Principle Study Design

Study Objective: Study the Efficacy and Safety of DARE-BV1 in the Treatment of Bacterial Vaginosis (n = 30)

Day 7 - 14 Day 21 - 30 Day 1 Baseline Visit Test-of-Cure Visit Continued Clinical Response Visit · Single dose administered · Patients questioned regarding · Patients questioned regarding comfort level & re-examined experience & re-examined Tests Performed: Tests Performed: Tests Performed: · Physiological symptoms · Physiological symptoms · Physiological symptoms • pH • pH Hq • · Saline "wet mount" · Saline "wet mount" · Saline "wet mount" • 10% KOH "whiff test" • 10% KOH "whiff test" · 10% KOH "whiff test" Urine pregnancy (if needed) Urine pregnancy (if needed) · Urine pregnancy (if needed)

- Eligibility: Female subjects 18 years or older with confirmed clinical diagnosis of BV
- · Primary Endpoint: Clinical Cure at Test-of-Cure visit (defined as resolution of clinical findings from baseline visit);
- Secondary Endpoints: Proportion of patients with therapeutic and bacteriologic cures,^{1,2}
- · Safety: Patients were questioned about their comfort level and adverse reactions they experienced.



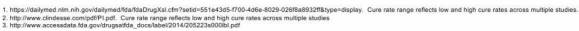
1. Therapeutic cure was a composite endpoint, which required both clinical cure (defined as clinical cure: resolution of all 4 Amsel criteria) and bacteriologic cure (Nugent score < 4). Bacteriologic cure required a Nugent score < 4. 2. Amsel & Gram Stain Criteria: https://www.cdc.gov/std/tg2015/bv.htm

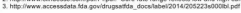
Bacterial Vaginosis DARE-BV1 Proof of Principle Study Design

A single dose of DARE-BV1 demonstrated high clinical cure rate compared to other approved products

Product	Clinical (Amsel) Cure	Bacteriologic (Nugent) Cure	Therapeutic Cure	
DARE-BV1 novel gel (clindamycin)	88%	57%*	57%*	
Solesec® ¹ (secnidazole 2g oral granules)	53-68%	40-46%	35-40%	
Clindesse®2 clindamycin phosphate Vaginal Cream, 2%	41-64%	45-57%	30-42%	
Metrogel, 1.3% 3	37%	20%	17%	

- * Based on data from 9 evaluable patients
- · 26 of 30 women completed the study
- Test-of-Cure Visit (Day 7 14)
 - · 23 of 26 (88%) women achieved clinical cure based on Amsel criteria
 - · 4 of 7 (57%) women had bacteriologic cure and 4 of 7 (57%) had therapeutic cure
- Continued clinical response visit (Day 21 30)
 - · 23 of 24 (96%) women showed continued clinical cure
 - · 8 of 9 women had bacteriologic cure and 7 of 9 had therapeutic cure





Perilgo



DARE-BV1

Bacterial Vaginosis Opportunity Overview

DARE-BV1 offers an attractive value proposition based on our belief that it has a low clinical risk profile, low development and regulatory costs, and an attractive commercial opportunity

Proof of principal study complete

- o 88% clinical cure rate in evaluable subjects
- o favorable efficacy profile compared to currently approved treatments

Same API (clindamycin phosphate 2%) as in currently approved treatment

Low Clinical Risk

Single Phase 3 clinical trial for FDA approval 7 Exploiting the 505(b)2 regulatory pathway

Low Development Cost
Anticipate less than \$10 million
Includes manufacturing, clinical trial, regulatory filing & action)

Approximately 21 M cases of BV reported annually in the US Approved prescription drugs have low patient share due to limited efficacy

Significant Market Opportunity



1. Based on prior sponsor communications with the FDA, one successful Phase 3 study with sufficient power and size may be sufficient for marketing approval in the U.S.



1. Global Market Insights, https://globenewswire.com/news-release/2016/05/19/841462/0/en/Contraceptives-Market-size-to-exceed-33-Billion-by-2023-Global-Market-Insights-Inc.html

Successful Proof of Concept Study

- Ovaprene demonstrated effectiveness in preventing sperm from entering the cervical canal in a proof-of-concept study in women (n=20):1
 - · No viable sperm in the cervical mucus
 - · No colposcopic abnormalities

CDRH (Device) Regulatory Pathway

· Single pivotal clinical trial expected for FDA approval

Attractive Market Opportunity

- >\$6 billion in US Rx sales of contraceptive products (2016).²
- 40 million women of reproductive age currently use a contraceptive method.³

Patent Coverage

- Patents covering the licensed technology have been granted with terms through 3Q 2028
- · Opportunity for Patent Term Extension (PTE) and potential new patents



Journal of Reproductive Medicine 2009; 54: 685-690
 IMS NSP through Dec 2016
 www.guttmacher.org, contraceptive fact sheet

Innovation in Contraception

Advances in hormone products have largely focused on reducing the hormone dosage, adjusting or extending the duration of protection and optimizing methods of administration.



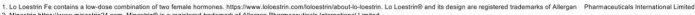






Convenience is driving new innovation

- NuvaRing®
 - · Monthly, convenient vaginal ring product form.
 - 2017 worldwide sales: \$761 million (Merck)⁵
- Mirena® Product Family
 - · Physician inserted, long-acting.
 - · Low/locally delivered hormone IUS.
 - 2017 worldwide sales: \$1.12 billion (Bayer)⁷



2. Minastrin https://www.minastrin24.com. Minastrin® is a registered trademark of Allergan Pharmaceuticals International Limited.
3. https://www.mirana-us.com/about-mirena/

5. Annual Report on Form 10-K for fiscal year ended December 31, 2017 6. Bayer Annual Report 2017. Includes sales for Mirena®, Kyleena® and Jaydess® / Skyla®



Women's Preferences

- 1. Effective Pregnancy Prevention
- 2. Convenient Product Forms
 - Independent surveys revealed that the vaginal ring has many of the features women deemed extremely important.1
- 3. Method Mix
 - >70% of women who practice contraception currently use non-coital (not in the moment) methods.2
- 4. Less Hormones
 - · A majority of women prefer a monthly option with a lower hormone dose than the pill.3

CONTRACEPTIVE METHOD CHOICE

Most effective method used in the past month by U.S. women, 2014

METHOD	No. of women	% of women aged 15-44	% of women at risk of unintended pregnancy	cont
Pill	9,572,477	15.6	22.7	
Tubal (female) sterilization	8,225,149	13.4	19.5	
Male condom	5,496,905	8.9	13.0	
ND D	4,452,344	7.2	10.6	
Vasectomy (male sterilization)	2,441,043	4.0	5.8	
Withdrawal	3,042.724	5.0	72	
Injectable	1,481,902	2.4	3.5	
Vaginal ring	905,896	15	21	
Fertility awareness- based methods Implant	832,216 965,539	1.3 1.6	20 23	
Patch	69,106	0.1	0.2	
Emergency contraception Other methods*	69.967 234,959	0.1	0.2	
No method, at risk of unintended pregnancy	4,408,474	72	10.5	
No method, not at risk	19,302,067	31.4	na	
Total	61,491,766	100.0	100.0	1 3

www.guttmac



Lessard, L.Perspectives on Sexual and Reproductive Health, Volume 44, Number 3,9-2012
 https://www.gutmacher.org/fact-sheet/contraceptive-use-united-states
 Hooper, DJ, Clin Drug Investig. 2010;30(11):74963

What's Missing in Contraception? Hormone free alternatives that are effective and easy to use

Least Effective			Hormone Free Product Landscape Marketed or in development		
100% Effective = 0%	Risk of Pre	gnancy	Spermicides / Vaginal Gels © Effectiveness (72% Typical Use)		
Birth Control Effectivenes	ss		Woman controlled		
Method	Perfect Use	Typical Use	Used "in the moment"		
Spermicide* / Vaginal Gels	82,00%	72.00%			
Sponge-Parous*	80.00%	76.00%	Condoms		
Sponge-Nulliparous*	91.00%	88.00%	Effectiveness (939/ Tunical Hea)		
Condom (male)*	98.00%	82.00%	Effectiveness (82% Typical Use)		
Diaphragm*	94.00%	88.00%	Not woman controlled		
Combined Pill & Progestin only*	99.70%	91.00%	A 11-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1		
Evra Patch*	99.70%	91.00%	Used "in the moment"		
Nova Ring*	99.70%	91.00%	Diaphragms		
Depo-Provera*	99.80%	94.00%			
IUD- ParaGard (Copper T)*	99.40%	99.80%	Effectiveness (88% Typical Use)		
IUD- Mirena (LNg)*	99.80%	99.80%	Moman controlled		
Implanon*	99.95%	99.95%			
Female Sterilization*	99-50%	99.50%	Used "in the moment"		
Male Sterilization*	99.90%	98.85%	Long-acting IUD		
			Long-acting IOD		
	1		Effectiveness (99% Typical Use)		
			Not woman controlled		
Most Ef	fective		Physician inserted		



^{1.} Trussell J. Contraceptive Efficacy. In Hatcher RA, Trussell J, Nelson AL, Cates W, Kowal D, Policar M. Contraceptive Technology: Twentieth Revised Edition. New York, NY: Ardent Media, 2011. 2. http://www.contraceptivetechnology.org/wp-content/uploads/2013/09/CTFailureTable.pdf

Ovaprene® Non-hormonal, Monthly Vaginal Ring

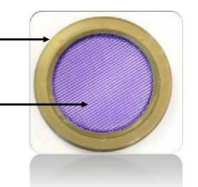
Spermiostatic Environment¹

- · Achieved through a contraceptive-loaded silicone ring matrix.
- · Releasing non-hormonal active Ferrous gluconate.

Physical Barrier1-

• 3-D, non-braided, fluid-permeable mesh barrier.

Rx distribution (OB/GYN) – anticipated upon approval.





New Contraceptive Option

Ovaprene® Overview

Ovaprene successfully prevented sperm from reaching the cervical canal in a previous human postcoital test (PCT) clinical study.

- 2009 Postcoital Assessment:
 - · Open-label, single-arm, pilot safety and tolerability study.
 - Published in the Journal of Reproductive Medicine, 2009.
- Patients:
 - N= 20; all women completed one cycle of use.
- - Postcoital testing revealed no viable sperm in the cervical mucus.
 - No colposcopic abnormalities, no significant changes in vaginal flora and no serious adverse effects observed.

Birth Control Effectivenes	s	
Method	Perfect Use	Typical U
Spermicide* / vugital gals	82.00%	72.00%
Sponge-Parous*	80.00%	76.00%
Sponge-Nulliparous*	91.00%	88.00%
Condom (male)*	98.00%	82.00%
Olaphragm*	94.00%	88.00%
Combined Pill & Progestin only*	99.70%	91.00%
Evra Patch*	99.70%	91.00%
Nova Ring*	99.70%	91.00%
Depo-Provera*	99.80%	94.00%
IUD- ParaGard (Copper T)*	99.40%	99.80%
IUD-Mirena (LNg)*	99.80%	99.80%
Implanon*	99.95%	99.95%
Female Sterifization*	99.50%	59.50%
Male Sterilization*	99.90%	58,85%



1. Journal of Reproductive Medicine 2009; 54: 685-690
2. Trussell J. Contraceptive Efficacy. In Hatcher RA, Trussell J, Nelson AL, Cates W, Kowal D, Policar M. Contraceptive Technology: Twentieth Revised Edition. New York, NY: Ardent Media, 2011.
3. http://www.contraceptivetechnology.org/wp-content/uploads/2013/09/CTFailureTable.pdf



New Contraceptive Option

Ovaprene® Overview

U.S. Regulatory Strategy

- · PMA with CDRH (Medical Device Division) as lead review division.
- · Pathway expected to be based on similar CDRH approvals Example: Caya® diaphragm.*

Step 1 - Postcoital test (PCT) 2018 / 2019*

- The study is enrolling 50 couples.
 - · 25 women complete a total of 21 visits
- · Evaluated over the course of five menstrual cycles.
- Each woman's cervical mucus will be examined at several points during the study:
 - · Cycle 1 Baseline (excludes the use of any product),
 - · Cycle 2 Use of a barrier method (diaphragm),
 - · Cycles 3,4 and 5 Ovaprene vaginal ring.
- Assess motile sperm per high powered field (HPF) in the cervical mucus, post coitus.
- · Safety assessments, PK, acceptability, fit, and ease of use.
 - · Data from the study is expected to be available in the second half of 2019.
 - If there is demonstration of feasibility in the PCT clinical trial, the Company intends to prepare and file an Investigational Device Exemption (IDE) with the FDA to commence a pivotal clinical trial to support marketing approvals of Ovaprene in the United States, Europe and other countries worldwide.

→ Step 2 – Pivotal Study 2020 / 2021*

- · Single pivotal clinical (expected).
 - N= ~250 completers over 6 months of use.
 - · Primary Endpoints: Safety & Efficacy
 - Pregnancy probability.
 - · Secondary Endpoints:
 - · Acceptability/product fit/ease of us
 - · Assessments of vaginal health.

ф daré

*Anticipated regulatory pathway and timelines. Daré has not had any communications with the FDA regarding the specific PMA requirements for Ovaprene.

Features Desired Most in Birth Control:14	Design Features of Ovaprene: ^{5,6}
Convenience (Easy to Use & Easy to Remember)	Monthly Ring Product Form Women chose rings for the convenience of a non-daily option.
1 Hormone Free	No Hormones in the API Unique dual action MOA (spermiostatic & barrier).
€ Efficacy	Expected Typical Use Effectiveness Comparable to Hormone Contraception (88% vs 91%).
Tavorable Side Effect Profile	No Colposcopic Abnormalities No significant changes in vaginal flora. No serious adverse effects observed in prior published study.
Easily Manage Fertility	No Systemic Activity Inserted and removed without a provider. Immediate return to fertility.



ф daré

https://www.urban.org/urban-wire/women-want-effective-birth-control
Lessard, L,Perspectives on Sexual and Reproductive Health, Volume 44, Number 3,9-2012
Hooper, DJ, Clin Drug Investig, 2010;30(11):74963
Ersek, J, Matern Child Health J (2011) 15:497-506
Journal of Reproductive Medicine 2009; 54: 685-690
Trussell J. Contraceptive Efficacy. In Hatcher RA, Trussell J, Nelson AL, Cates W, Kowal D, Policar M. Contraceptive Technology: Twentieth Revised Edition. New York, NY: Ardent Media, 2011.



1. https://www.visiongain.com/sexual-dysfunction-drugs-market-will-reach-7-7bn-in-2019-predicts-a-new-visiongain-study/

Female Sexual Arousal Disorder (FSAD) Sildenafil Cream, 3.6%

Topical Cream Sildenafil Cream, 3.6% Top line

Successful Proof of Concept

- Sildenafil Cream, 3.6% improved genital blood flow in a proof-of-concept study (n=31):1
 - · Efficacy signal observed in both pre and postmenopausal patients
 - · Excellent systemic/local safety and tolerability profile

505(b)(2) Regulatory Pathway

· Ability to leverage the safety profile of sildenafil (Viagra®) for FDA submission package

Attractive Market Opportunity²

- · 33% of females in the U.S. (21 to 60 years old) experience symptoms of low or no sexual arousal
- 16% (~10m women) are considered distressed and are seeking a solution to improve their condition

Patent Coverage

- Patents covering the licensed technology have been granted with terms through 2031 (through June 2029 in the U.S.)
- · No ANDA route: ANDA is not currently an option for topicals that result in low systemic uptake



1. Data on file

2. Ad Hoc Market Research: FSAD Prevalence Report (Oct 2015) conducted for SST LLC. Based on US Census projections for 2016

Female Sexual Arousal Disorder (FSAD)

Sildenafil Cream 3.6%

Dyspareunia

Vulvar-Vaginal Atrophy

Hypoactive Sexual Desire Disorder (HSDD)

Female Sexual Arousal Disorder (FSAD)













With its approval of Addyi®, FDA has now acknowledged and formally classified the distinct and separate disorders that comprise Female Sexual Dysfunction.

Where HSDD is characterized primarily by a lack of sexual desire, FSAD is characterized primarily by an inability to attain or maintain sufficient physical sexual arousal.

- INTRAROSA is a registered trademark of Endoceutics, Inc.

- Int tarkUSA is a registered trademark of Endoceutics, inc.

 · Imwexxy is a trademark of TherapeuticsMD, Inc.

 · Osphena is a registered trademark of Duchesnay USA, Pennsylvania, USA, ESTRACE® is a registered trademark of Allergan Pharmaceuticals Internatic

 Premarin is a registered trademark of Pfizer Inc.

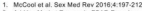
 Addyl is a registered trademark of Sprout Pharmaceuticals, Inc.



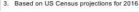
Female Sexual Arousal Disorder (FSAD) Sildenafil Cream 3.6%

FSAD is characterized primarily by an inability to attain or maintain sufficient physical sexual arousal; it is also characterized by distress or interpersonal difficulty.*

- Estimated 23-33% of women suffer from arousal disorder:
 - Meta-analysis of 95 studies from 2000-2014 indicated the prevalence of Female Sexual Dysfunction in premenopausal women worldwide is 40.9%, and difficulty with arousal alone is 23%.1
 - 33% of women in the U.S. age 21 to 60 (approximately 20 million women), experience symptoms of low or no sexual arousal.2,3
 - 10 million women are considered distressed and actively seeking treatment.2



McCool et al. Sex Med Rev 2016;4:197-212.
 Ad Hoc Market Research: FSAD Prevalence Report (Oct 2015) conducted for SST LLC.
 Based on US Census projections for 2016.





Female Sexual Arousal Disorder (FSAD) Sildenafil Cream 3.6%

Key Takeaways of Viagra® studies:

- · Oral sildenafil (Viagra) demonstrated statistically significant ac
- Side effects of the oral formulation led to the investigation of a topical route of administration

Increased blood flow and clinical efficacy with oral sildenafil (Viagra®) in women:

- Statistically significant increases in Vaginal Pulse Amplitude (VPA)
- Statistically significant improvement in genital stimulation (FIEI)2

Pfizer Clinical Field Study - Oral Viagra Pfizer VPA Clinical Lab Study - Oral Viagra Improvement on FIEI Mean and Maximum VPA† Questions† Vaginal Pulse Amplitude (mV) 6 Observed Number Improved (%) 60 50 4 40 3 30 20 Question 2 Question 4 Mean (Erotic) Maximum (Erotic) ■ Placebo ■ Oral Viagra®

† Twelve healthy premenopausal women were studied.

■ Placebo ■ Oral Viagra®

Female Intervention Efficacy Index (FIEI)

estion #2 – "After taking study medication, the sensation/feeling in my genital (vaginal, labia, clitoris) area during intercourse or stimulation (foreplay) med to be: (a) more than before, (b) less than before, or (c) unchanged". Question #4 – "After taking the study medication, intercourse and/or foreplay was: pleasant and satisfying, better than before taking the study medication, (c) unchanged; no errence, or (d) pleasant; but still not like it used to be or I vouid like it to be." 202 potemporausal women with FSAD who had protocol specified estradiol free testosterone concentrations, and/or were receiving estrogen and/or androgen replacement therapy were studied.



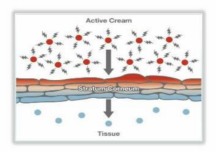
1. The Enhancement of Vaginal Vasocongestion by Sildenafil in Healthy Premenopausal Women. Journal of Women's Health & Gender-Based Medicine. Vol. 11, No. 4. 2002 2. Safety and Efficacy of Sildenafil Citrate for the Treatment of FSAD: A Double-Blind, Placebo Controlled Study. The Journal of Urology. Vol 170, 2333-2338, December 2003.

Female Sexual Arousal Disorder (FSAD)

Sildenafil Cream 3.6%

Formulation Innovation

- Sildenafil Cream, 3.6% designed to directly increase local blood flow to the genital tissue.
- The formulation delivers localized action, with minimal systemic uptake of the active drug.¹



SST Formulation Technology

6 issued patents in the U.S. on the topical delivery of Sildenafil and other PDE-5 inhibitors.

- Leveraging the known therapeutic benefit of oral sildenafil to stimulate increased blood flow to the genital tissue.
- If approved, Sildenafil Cream, 3.6% may offer a safe, effective and 'on demand' solution to difficulties with sexual arousal.



1. Data on file

Female Sexual Arousal Disorder (FSAD)

Sildenafil Cream, 3.6%

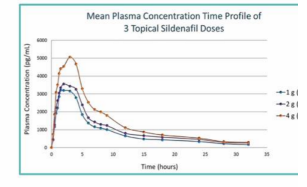
Phase 1 Study of SST-6007 (Sildenafil Cream, 3.6%)1

- Normal healthy postmenopausal women were dosed with escalating doses of Sildenafil Cream, 3.6%, using a cross-over study design.
- Topical sildenafil had significantly lower systemic exposure compared to a 50 mg oral sildenafil dose
 - AUC 3-6%
 - C_{max} 1-2%
- Safe and very well tolerated at clinically relevant doses (1-2g)
- Favorable product characteristics as self reported by subjects
 - · Easy to use
 - Readily absorbed

Phase 2a Study of SST-6007 (Sildenafil Cream, 3.6%)1

 Demonstrated increased blood flow in the genital tissue compared to placebo (mean change in VPA analysis) in 31 women (pre and postmenopausal) ~30 minutes post dosing

Treatment	N	Sildenafil Single Dose	C _{max} (ng/ml)	T _{max} (hr)	AUC (h*ng
Topical Sildenafil 1 g of cream	20	35 mg	3.4	2.37	25
Topical Sildenafil 2 g of cream	20	71 mg	3.8	2.27	30
Topical Sildenafil 4 g of cream	19	142 mg	5.3	2.22	42





Female Sexual Arousal Disorder (FSAD)

Sildenafil Cream 3.6%

Phase 2b Program: Continue to explore additional clinical and non-clinical work that might be valuable or required to support the overall program and the anticipated design of the Phase 2b.

Content Validity

A content validity study is designed to help ensure

- the concepts we plan to measure are the most important and relevant to our target population.
- · This is a non-interventional study participants will not be asked to use or evaluate any products.

Planned Type C Meeting

 We will request at Type C meeting to get feedback on whether the agency agrees that the patient (PRO) reported outcomes instruments are content valid for the target population.

At Home Study

- The Phase 2b at-home study will allow patients to use investigational product and placebo in their home setting.
- The FDA is agreeable to a 12-week Phase 2b for Sildenafil Cream, 3.6% to assess reasonable safety and preliminary efficacy.
- The 2016 Draft Guidance reflects expectations regarding Phase 3 study length and patient population.

Key Takeaways:

The Phase 2b program will consist of a content validation component (ongoing), followed by at-home dosing of the investigational product and a placebo control. The plan is to use the selected PRO instrument and FDA agreed upon endpoints for the Phase 2b and Phase 3 clinical trials.



Female Sexual Arousal Disorder (FSAD) Sildenafil Cream 3.6%

Dyspareunia

Vulvar-Vaginal Atrophy

Hypoactive Sexual Desire Disorder (HSDD)

Female Sexual Arousal Disorder (FSAD)













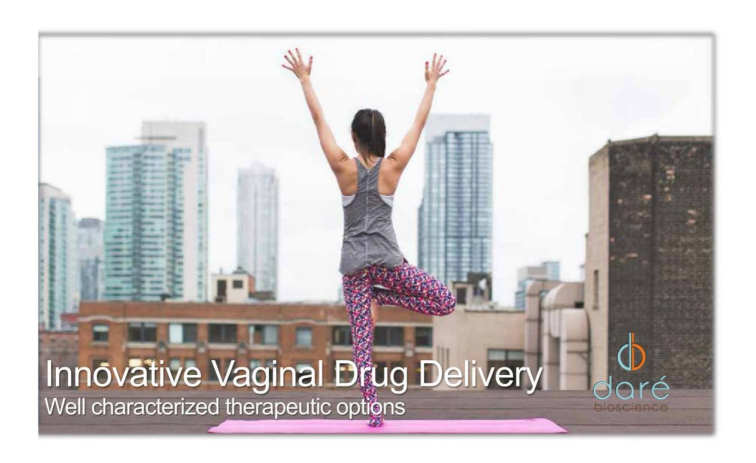
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 Premarin is a registered trademark of Pfizer Inc.
 Addyl is a registered trademark of Sprout Pharmaceuticals, Inc.





Intravaginal Ring (IVR) Technology Platform

Daré has an exclusive, global license to Juniper's novel IVR technology originally developed by Dr. Robert Langer from MIT¹ and Dr. William Crowley² from Massachusetts General Hospital and Harvard Medical School. Daré's exclusive license covers all rings in development as well as additional applications of the IVR technology platform in other therapeutic areas.

- · Features of the Juniper intravaginal ring technology include:
 - · Sustained drug delivery
 - · Variable dosing and duration
 - Single or multiple drug delivery via a solid ethylene vinyl acetate polymer matrix (without the need for a membrane or reservoir to contain the active drug or control the release)
- Current 505(b)(2) candidates licensed from Juniper include:

DARE-HRT1

 Formerly JNP-0201, a combination bio-identical estradiol + bio-identical progesterone ring for hormone replacement therapy

DARE-FRT1

 Formerly JNP-0301, a bio-identical progesterone ring for the prevention of preterm birth and for fertility support as part of an IVF treatment plan



Hormone Replacement Therapy (HRT)

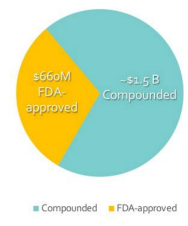
DARE-HRT1 (bio-identical estradiol + progesterone)



HRT remains the most effective treatment for vasomotor symptoms (VMS) and the genitourinary syndrome of menopause (GSM) and has been shown to prevent bone loss and fracture.1

- 45M women in U.S. approaching or in menopause.²
- 2012 NAMS consensus statement supports HRT in peri- and postmenopausal women – estrogen to reduce symptoms and progesterone to prevent thickening of uterine wall.3
- NAMS recommends non-oral route over oral.3
- · 2002 Women's Health Initiative (WHI) study showed that the long-term use of certain synthetic hormones (a combination of medroxyprogesterone and conjugated equine estrogens) increased the risk of breast cancer, stroke, heart attack and blood clots







^{1.} The 2017 hormone therapy position statement of The North American Menopause Society; Menopause: The Journal of The North American Menopause Society Vol. 24, No. 7, pp. 728-753 2. U.S. Census Bureau, Population Division. Table 2. 2015 to 2060 (NP2012-T2). Released Dec. 2012. 3. Menopause, Vol. 19, No. 3, 2012.

4. U.S. 2014. Source: Symphony Health Solutions Report

Hormone Replacement Therapy (HRT)

DARE-HRT1 (bio-identical estradiol + bio-identical progesterone)

Phase 1 - HRT

DARE-HRT1 for the treatment of VMS due to menopause – combination bio-identical estradiol and progesterone in a convenient 28 day IVR

- · Proposed Study:
 - A Phase 1, Open-Label, 3-arm Parallel Group Study to Evaluate the Pharmacokinetics and Safety of DARE-HRT1 (80 μg and 160 μg Estradiol/ 4 mg and 8 mg Progesterone Intravaginal Rings) in Healthy Post-Menopausal Women.
- Primary Objectives:
 - To describe the PK parameters over 28 days using two different dose combinations of DARE-HRT1 Intravaginal ring (IVR):
 - · Estradiol 80 μg/Progesterone 4 mg IVR
 - Estradiol 160 µg/Progesterone 8 mg IVR
 - Identify the steady-state PK after 28 days of each DARE-HRT1
- N=30



Pregnancy Maintenance

DARE-FRT1 (bio-identical progesterone)

- Prevention of Pre-term Birth (PTB)
 - In the US, approximately 12% of pregnancies are preterm (less than 37 weeks)¹
 - Standard interventions include steroids, hormones and tocolytic agents to stop/slow the frequency and duration of contractions2
- Assisted Reproductive Technologies (ART)
 - The global ART market is expected to reach USD 45 billion by 2025, according to a new report by Grand View Research, Inc.3
 - · Childbearing postponement is a high impact driver of the infertility treatment market.
 - Increasing marital age, rising tobacco and alcohol consumption, and increasing obesity rates are some of the other factors contributing to the market growth.
 - Furthermore, increasing incidence rate of conditions such as poly-cystic ovarian syndrome (PCOS), tubal factors and endometriosis are other drivers of the market.
 - Record number of women using IVF to get pregnant ⁴
 - More American women have had medical help to have their babies than ever, according to the latest annual report from the Society for Assisted Reproductive Technology.



https://www.stanfordchildrens.org/en/lopic/default?id=prematurity-90-P02401 https://www.uptodate.com/contents/preterm-labor-beyond-the-basics https://www.grandviewresearch.com/press-release/jobal-assisted-reproductive-technology-ma https://www.cnn.com/2014/02/17/health/record-ivf-use/index.html

Vaginally Delivered Tamoxifen for VVA DARE-VVA1

Vaginally Delivered Tamoxifen to treat VVA in HR+ Breast Cancer Patients

- DARE-VVA1 (Formerly PT-101)
 - · A proprietary formulation of tamoxifen for vaginal administration.
 - · Potential to be the first treatment specifically approved for the treatment of vulvar and vaginal atrophy (VVA) in patients with hormone-receptor positive (HR+) breast cancer.
- VVA is a chronic condition characterized by pain during intercourse, vaginal dryness and irritation.
 - · Most women use localized estrogen therapy which is contraindicated for the more than two million women diagnosed with, or at risk of recurrence of, ER-positive and PR-positive breast cancer.1
 - · Daré intends to develop this novel local application of tamoxifen to mitigate the symptoms of VVA for patients with or at risk for hormone-receptor-positive breast cancer, including women currently on anticancer therapy.
 - Due to the use of aromatase inhibitors for the treatment of HR+ breast cancer, the prevalence of VVA in postmenopausal breast cancer patients is reported to be between 42 and 70 percent.²



Vaginal Tamoxifen – Proof of Concept Study¹

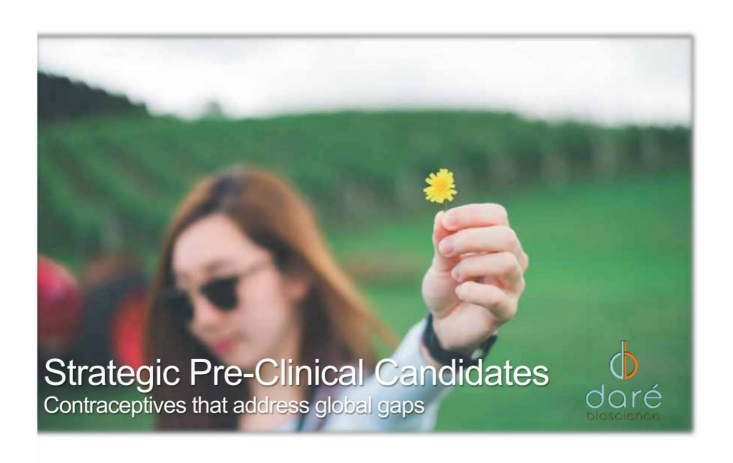
· This exploratory study in four postmenopausal women diagnosed with VVA demonstrated that a selfadministered vaginal suppository containing tamoxifen (20mg) dosed daily for one week and twice weekly for three months was effective in reducing vaginal pH and vaginal dryness

Vaginal Tamoxifen	Enrollment (Baseline)	On Treatment (Month 3)	Paired Difference (Baseline vs. Month 3)
Median Vaginal pH Lower pH value is a measure of symptom relief Normal vaginal pH is usually less than 4.5.2	7.1 range 6.5 to 7.5	5.0 range 5.0 to 5.2	-2.0 median range -2.5 to -1.5
Vaginal Dryness Rated using a visual analogue scale (VAS) that ranged from: 0 = Not bothered by dryness 10 = Extremely bothered by dryness Decreased vaginal dryness is a measure of symptom relief	8.0 range of 7.5 to 9.0	3.0 range 2.0 to 3.0	-5.5 median range -6.0 to -4.5

- In addition, systemic absorption of tamoxifen was not significant.
 - After 8 weeks of study treatment with vaginal tamoxifen, the median plasma concentration of tamoxifen was 5.8 ng/ml, with a range of 1.0 to 10.0 ng/ml
 - In comparison, after 3 months of administration of 20mg, once-daily oral tamoxifen citrate (Nolvadex),2 the average steady st plasma concentration of tamoxifen is 122 ng/ml with a range of 71 to 183 ng/ml



1. Clin. Exp. Obstet. Gynecol. - ISSN: 0390-6663 XLVI, n. 2, 2019
2. https://www.medicalnewstoday.com/articles/322537.php
3. US Food and Drug Administration: "Drug Approval Package: Nolvadex (Tamoxifen Citrate) NDA# 21-109.2002". Available at: https://www.accessdata.fda.gov/drugsatfda_docs/nda/2002/21109_Nolvadex.cfm



A New Long Acting Contraceptive Option

Injectable Etonogestrel ORB 204 & 214

Microparticle 6 & 12 Month Injectable Contraception

ORB-204 and ORB-214, injectable etonogestrel¹

The initial development on Orbis' long-acting injectable contraceptive program was carried out under a subcontract funded by Family Health International (FHI 360) through a grant from the **Bill & Melinda Gates Foundation**.

- · Pre-clinical studies for the 6- and 12- month formulations have been completed to date:
 - · Establishing pharmacokinetics and pharmacodynamics profiles.

An injectable contraceptive is designed to provide discreet, non-invasive protection over several months

• Limitations of the currently marketed injectable contraceptive: provides contraceptive protection for only three months, and can delay the ability to get pregnant for up to ten months after receiving the injection.

Target product profile of long-acting injectable

 Prolonged duration (6 to 12 months), improved ease of use, with an improved side effect profile and predictable return to fertility.



Data on file

A New Contraceptive Target DARE-RH1 CatSper

Ca2+ Target DARE-RH1

A Novel Approach To Male And Female Contraception.

- The identification of the CatSper target represents the potential to develop a novel class of non-hormonal contraceptive products for both men and women.
 - The discovery of a sperm-specific ion channel, CatSper, was validated in animal models where it was demonstrated that male mice lacking CatSper have poor sperm motility.
- CatSper proteins are ion channels expressed solely in the membranes of sperm flagellum and are essential to sperm motility.
- Pre-clinical research has demonstrated CatSper mediates hyperactive motility of sperm.
 - Sperm hyperactivity is necessary to penetrate the physical barrier known as the zona pellucida which encloses the ovum and protects the egg.¹
 - The contraceptive benefit of targeting CatSper is achieved by inhibiting sperm hyperactivity and preventing egg fertilization.



1. http://journals.plos.org/plosone/article?id=10.1371/journal.pone.0028359



Investment Highlights

Financial Profile

Background

- NASDAQ:DARE
- Publicly traded via reverse merger that closed July 19, 2017

Balance sheet, December 31, 2018:

- · \$6.8 million in cash
- · Non-dilutive NIH SBIR Award:
 - In Q1-2019 Daré received a second Notice of Award of \$982,851, part of a grant providing up to \$1.9 million in the aggregate for Ovaprene® research from the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), a division of the National Institutes of Health (NIH). The first award under the grant in 2018 was \$224,665.
 - This second Notice of Award, for the additional \$982,851, followed the NIH's review of a data analysis and other results of the first phase of work.
- 11.4 million common shares and 3.75 million warrants outstanding
- No debt



Management Team Daré Bioscience

Sabrina Martucci Johnson, MSc, MIM President and CEO	Cypress Bioscience, Calibr, Advanced Tissue Sciences, WCG, Baxter Healthcare
Lisa Walters-Hoffert Chief Financial Officer	ROTH Capital Partners, Citicorp Securities, Bank of America, Oppenheimer & Co.
David Friend, PhD Chief Scientific Officer	Evofem, CONRAD, Elan Corporation
John Fair Chief Business Officer	Evofem, WCG, Gemini Healthcare, Aegis plc
Mark Walters Vice President, Operations	Pacira, SkyePharma, Alliance Pharmaceuticals, American Home Products
Mary Jarosz, RPh, RAC, FTOPRA Global Head of Regulatory Affairs	Evofem, WCG, Abbott Laboratories
Christine Mauck, MD, MPH Medical Director	CONRAD, Population Council, RW Johnson, FDA
Bridget Martell, MD, MA Medical Affairs	Juniper Pharmaceuticals, Purdue Pharma, Pfizer
Nadene Zack, MSc Sr. Director Clinical Operations	Retrophin, Aragon, Cypress Bioscience, Pfizer



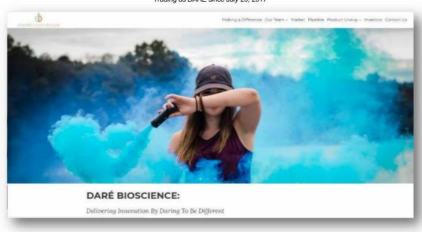
Board of Directors Daré Bioscience

Roger Hawley (Chairman)	Zogenix, Alios Biopharma, Cypress Bioscience, InterMune, Elan Corporation, GSK
Jessica Grossman, MD	Medicines360, Sense4Baby, Johnson & Johnson
Susan Kelley, MD	Bayer, BMS, ArQule, Cerulean
Greg Matz	CooperSurgical - Cooper Companies, Agilent, Hewlett Packard
William Rastetter, PhD	Neurocrine Biosciences, IDEC, GRAIL, Receptos, Illumina, Cerulean
Robin Steele, JD, LLM	InterMune, Elan Corporation, Alveo, Alios Biopharma
Sabrina Martucci Johnson, MSc, MIM	Cypress Bioscience, Calibr, Advanced Tissue Sciences, WCG, Baxter Healthcare



Corporate & Investor Communications

NASDAQ: DARE Trading as DARE since July 20, 2017







www.darebioscience.com







Bacterial Vaginosis Market Insights

American Sexual Health Association (ASHA), in conjunction with Harris Poll, conducted a national survey of 304 women ages 18 to 49 who have had bacterial vaginosis (BV). The survey was conducted online by Harris Poll on behalf of Symbiomix Therapeutics, LLC, a Lupin company, and the ASHA within the United States between September 14 and 29, 2017 among 304 US women aged 18-49 who have been diagnosed by a healthcare professional with BV within the past 2 years ("women with bacterial vaginosis").

Bacterial Vaginosis Market Insights American Sexual Health Association (ASHA) National Bacterial Vaginosis Survey

 <u>76%</u> of women with BV stated they would have gone to <u>see a healthcare professional sooner</u> if they were aware of the risks associated with BV if left untreated

IF BV RISK FACTORS WERE KNOWN

- Would Seek Treatment Sooner
- Would Not Seek Treatment Sooner



 Only 43% of women with BV are aware that if left untreated, BV can cause an increased risk of sexually transmitted infections (STIs)

AWARE OF LINK TO STI

- Aware BV Can Increase Risk of STI
- Unaware BV Can Increase the Risk of STI

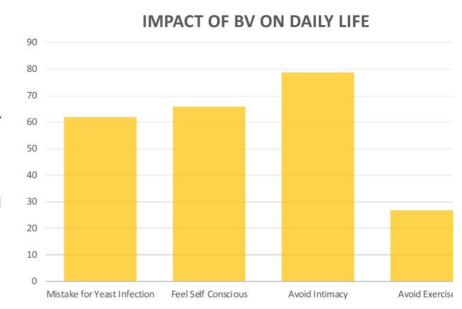




http://www.ashasexualhealth.org/understanding-womens-experiences-with-bacterial-vaginosis/

Bacterial Vaginosis Market Insights American Sexual Health Association (ASHA) National Bacterial Vaginosis Survey

- According to the ASHA survey, 62% of women mistake BV for a yeast infection prior to diagnosis
- Most women with BV feel self-conscious (68%) and/or embarrassed (66%) due to their condition
- Women with BV avoid everyday activities including being intimate with their spouse/partner (79%), working out (27%), or going on a first date (17%)





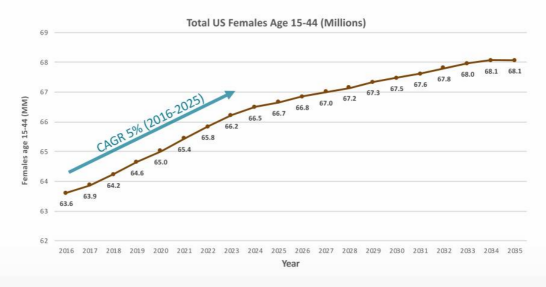
http://www.ashasexualhealth.org/understanding-womens-experiences-with-bacterial-vaginosis/



Ovaprene Market Insights

Secondary Market Research & Market Sizing Data Prepared by SmartPharma, February 2019. Data on File.

US Contraceptive Population is Over 60 million and Continues to Grow



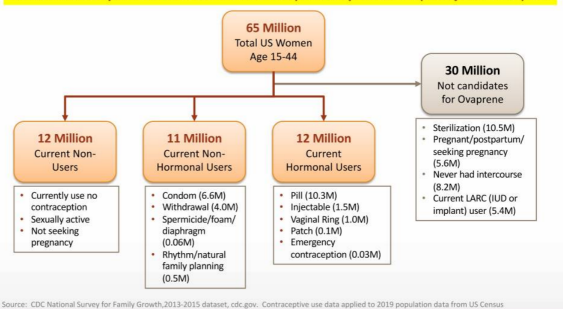
Source: US Census Bureau, 2017 National Dataset (2016 is base population estimate for projection) https://www.census.gov/programs-surveys/popproj.html

Dare Bioscience - Ovaprene Market Assessment, Jan 2019



Ovaprene Potential – Total Market Size

There are currently 35 million US women who could potentially choose Ovaprene for contraception



Dare Bioscience – Ovaprene Market Assessment, Jan 2019



Negative Information About Hormones is Persistent in the Public Domain

As a non-hormonal option, Ovaprene does not have to overcome myths or negative "press"

5 Reasons Women Avoid Birth Control

Reason #1: "I don't want to get fat"

Reason #2: "It might make me depressed" Reason #3: "Birth control causes cancer"

Reason #4: "I don't want to put chemicals in my body"

Reason #5: "I'm not at risk for getting pregnant"

6 Reasons Why You Shouldn't Take The Pill Long Term

April 4, 2017 by Fertility Friday / 21 Comments

- · The pill lowers your sex drive
- The pill shrinks your clitoris and causes painful sex
- · The pill causes depression and anxiety
- Long term pill use puts you at an increased risk of cervical cancer
- Long term pill use is associated with a delay in your return to fertility



1. Hormonal Birth Control Comes with Side Effects

- 2. Birth Control is Full of Hormones/Chemicals
- 3. Birth Control Works Against Your Body
- 4. Birth Control *May* Cause Abortions
- 5. A Whole Host of Other Reasons

HelloFlo, Mar 22, 2017, www.helloflo.com
Fertility Friday, April 4,2017, www.fertilityfriday.com
Equipping Godley Women, April 15, 2015, www.equippinggodleywomen.com

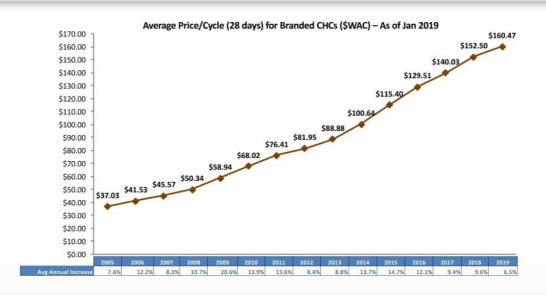


Contraceptive Pricing and Reimbursement

Dare Bioscience – Ovaprene Market Assessment, Jan 2019



Brand Contraceptives Have Consistently Increased in Price

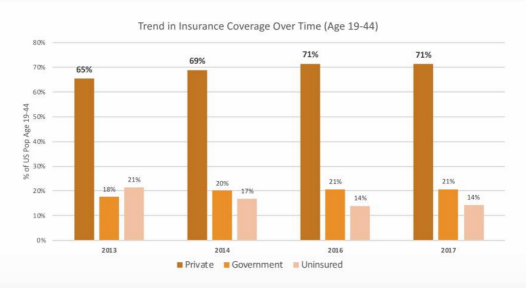


Source: MediSpan PriceRx, accessed Jan 2019. Average of 13 available branded contraceptives available in US market. Only three of these brands have no generic equivalent, and the average price/cycle for those 3 = \$171.06

Dare Bioscience - Ovaprene Market Assessment, Jan 2019



Over 70% of Reproductive-Aged Women in the US Have Private Insurance



*Patients can have more than one form of insurance, so totals may exceed 100%)

Source: Berchick et al. US Census Bureau, Health Insurance Coverage in the US: 2017, Issued Sep 2018

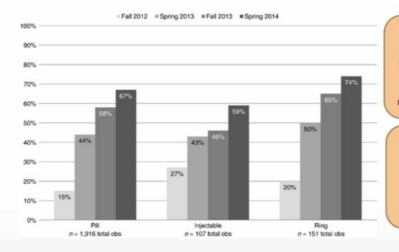
Dare Bioscience - Ovaprene Market Assessment, Jan 2019



Most Women Pay \$0 For Birth Control Since the ACA Was Enacted

Percent of Women with \$0 Copay for Birth Control Over Time

(n=892 women age 18-39 with private health insurance who used a prescription contraceptive method)



By spring 2014, mean and median out-of-pocket costs for the pill were \$6.48 and \$0 per month, respectively

HHS issued a clarification in May 2015 that required coverage of at least 1 product per method with \$0 copay – therefore the percentages have likely increased since this data

Sources+

Guttmacher Institute Continuity and Change in Contraceptive Use Study in: Sonfeld A, et al. Contraception 2015;91:44-48 US Dept of Health and Human Services (HHS) FAQs About ACA Implementation (Part XXVI), www.cms.gov

Dare Bioscience – Ovaprene Market Assessment, Jan 2019



ACA Contraceptive Mandate: Current Status of Contraceptive Policy

- Insurance plans must cover all FDAapproved methods with no copay or cost sharing to patients¹
 - They must cover at least one type of each method in each category
- 62.8 million women (age 18-64) now have birth control coverage with no cost sharing²
 - Exemptions and accommodations for religious and moral objections are in place, but they affect only 6,400 to 127,000 women³
 - The latest attempts by the current administration to broaden the exemption has been blocked by two federal courts⁴
- If Ovaprene is approved, it has the potential to be the only product in the category, as it is a vaginal ring with a spermiostatic active.

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-www.nealtricare.gov/	coverage/birth-control-bene	SHEST

²National Women's Law Center Fact Sheet, Nov 2018 ³HHS Fact Sheet, Nov 7 2018

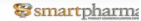
⁴National Women's Law Center Fact Sheet, Oct 2018 and Washington Post, Jan 14, 2019

Minimum Contrace	ptive Coverage Requirements Clar	ified by HHS Guidance
Contraceptive Method	Preducts/Options	Must Cover
Surgical sterilization	Also called tubal ligation	V
Implant sterilization	Only Essure available	V
Implantable Rod	Multiple	√acleast1
IUO - Copper	Only ParaGard available	4
IUD - Frogestin	Multiple	√at least 1
Injection	Multiple	√at least 1 (may be genesic)
Oral contraceptives - combined	Multiple	√at least 1 (may be generic)
Oral Contraceptives - progestinuosily	Multiple	√acleast 1 (may be generic)
Oral Contraceptives - extended/continuous use	Multiple	√at least 1 (may be generic)
Patch	Multiple**	√at least 1 (may be generic)
Vaginal Ring	Only Novelling available	✓
Diaphragm with Spermicide	Grily Miles Connilles available	4
Spange with Spermicide	Only Today Sponge available	4th
Cervical Cap with Spermicide	Only FemCap available	V
Female Condom	Multiglie	g/a
Spermicide alone	Multiple	√at least 1 (may be generic)
Emergency Contraception- Progestin	Multiple	√ at least 1 (may be generic)
Emergency Contraception Ulipristal Acetate	Only ella available	√

*Approved for sale own the counter but only covered at no cost with a prescription.

"The manufacturer of the brand name (Orthobina) patch has discontinued production and the generic atternative will be the only patch available.

IURCES: FDA, Birth Control Guide: and Depts of Labor, Health and Human Services, and Treasury, FAQs abo fordable Care Act Implementation (Part XXVII).





Sildenafil Cream, 3.6% FSAD Demographic Insights & Concept Test

Market Research Report Conducted by Ad Hoc Research on behalf of Strategic Science & Technologies, LLC. 222 Third Street, Suite 2242, Cambridge, MA 02142 – December 2015

FSAD - Psychological & Physiological Impact



The Current Experience of FSAD Sufferers

(Physical and Psychological)

Experience of FSAD Sufferers

- The concept definitely has potential. FSAD sufferers are likely to purchase it and are willing to give it a try.
- · A few questions remain:
 - 1. What do they currently experience during sexual activity that they are hoping the cream will rectify?

They often feel	They do not often feel
The inability to attain an adequate level of sexual excitement	Genital tenseness or tightness
The lack of desire for intimacy	Genital pulsing or throbbing
Lack of genital or clitoral fullness, pressure or engorgement	The feeling of muscle contractions in their genitals
Lack of genital wetness or lubrication	The feeling of readiness
	Satisfaction with their level of physical arousal

2. What are their main physical desires when it comes intimacy?

- They are desperate for their bodies to respond, be it to...
- · Intimacy,
- · An intimate touch;
- Touch





Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47.

Experience of Female Sexual Arousal Disorder (FSAD) Sufferers

3. What is the psychological impact of this disorder?

- The impact appears to be immense. Emotions run the gamut from dissatisfaction with to anger about their sex lives.
- The most frequent feelings include:
- · Dissatisfaction with their sex lives;
- Bothered by their low sexual desire;
- . Unhappiness about their sexual relationships; and
- · Frustration due to their sexual problems.
- Thus, conveying an understanding of these feelings, either in claims, in communications or both, will promote interest in the product.



4. What "remedies" have they tried to combat the disorder?

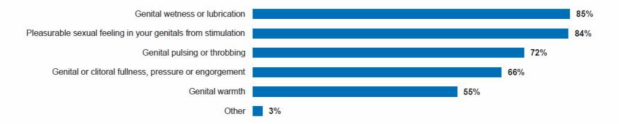


- Almost all FSAD sufferers surveyed have tried "something" to treat their difficulties getting or staying physically aroused.
- The most common are topical lubricants and a vibrator/other accessory for stimulation.



Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47.

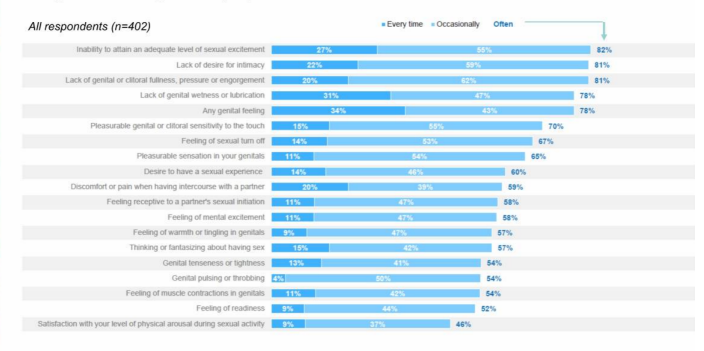
Female Sexual Arousal Disorder (FSAD) Respondents Indicators of Sexual Arousal



	Age group		FSAD LT SA 35-54 *	
All respondents (n=402)	21-44 n=195	45-60 n=207	Yes n=120	No n=282
Genital wetness or lubrication	87%	81%	83%	85%
Pleasurable sexual feeling in your genitals from stimulation	84%	84%	85%	84%
Genital pulsing or throbbing	75%	69%	69%	74%
Genital or clitoral fullness, pressure or engorgement	65%	68%	66%	66%
Genital warmth	58%	50%	51%	56%
Other	3%	2%	2%	3%

* LT- in a long-term relationship SA - currently sexually active 35-54 - ages of 35 to 54

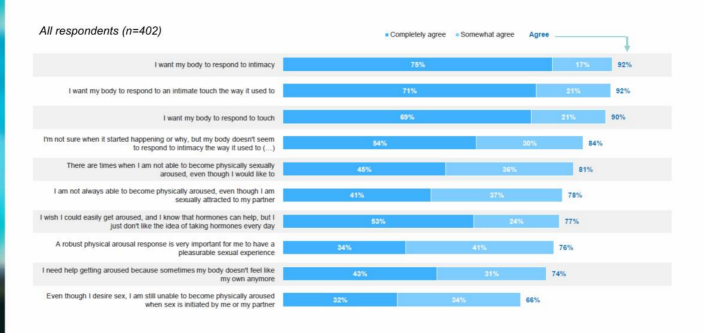
Female Sexual Arousal Disorder (FSAD) Respondents Signs & Symptoms



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Source data on File: Research Report – December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47.

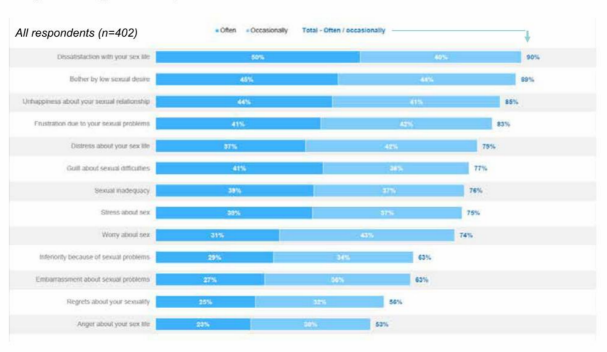
Psychological Impact of FSAD





Source data on File: Research Report – December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47.

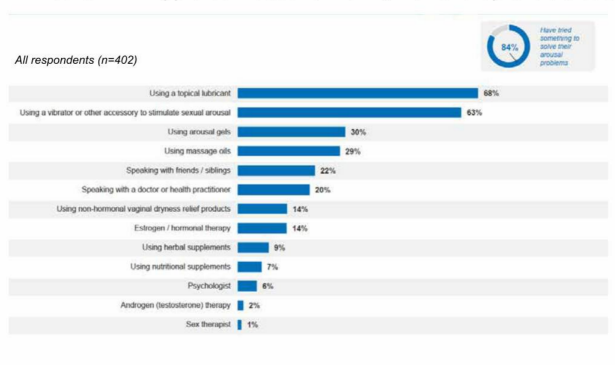
Psychological Impact of FSAD



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Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47.

Without an FDA Approved Product for FSAD, Women's Options are Suboptimal



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Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 41-47

Sildenafil Cream, 3.6% Product Profile Market Research



Sildenafil Cream, 3.6% Concept Test

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Sildenafil Cream, 3.6% Concept Testing



402 American women between the ages of 21 and 60 and suffering from Female Sexual Arousal Disorder (FSAD) were surveyed via web panel between November 18 and 23, 2015.



The purpose of this study is to measure the market potential of a topical cream version of sildenafil targeting women as a potential remedy for FSAD.



What degree of consumer interest is there in this product idea?

- A significant degree. Many FSAD sufferers have been experiencing symptoms for more than a year. In addition to the physical symptoms they experience, the psychological impact of the disorder is quite burdensome.
- On average, FSAD sufferers have tried approximately 3 different remedies chief among them topical lubricants and vibrators. However, they have found little to no relief.
- FSAD sufferers like the idea. They perceive it to be different from other remedies they have put to the test and they believe it will meet their needs.
- FSAD sufferers are ready to try something new especially one that promises no side effects.



What are the potential drivers of and barriers to adoption of the product?

Potential Drivers	Potential Barriers		
They want to give it a try. They are ready to try something new.	Embarrassment (in front of their partners, doctors and pharmacists).		
They want to increase their sex drive/sexual arousal.	Believability: will it work? They have tried many other "remedies" that have not.		



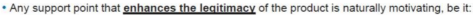
Source data on File: Research Report –December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 5

Sildenafil Cream, 3.6% Concept Testing



What are the most motivating claims?

- In concrete terms, the elements of the concept they like the most are:
 - √ No side effects (the #1 claim, by a very wide margin);
 - Proven safe;
 - ✓ Clinically tested;
 - ✓ Odorless;
 - Absorbs completely;
 - Available without a prescription.



- ✓ Doctor recommended;
- ✓ Available by prescription only for two years before being available without a prescription;
- ✓ The same active ingredient as in *Viagra* (although slightly less so than the previous two).



Source data on File: Research Report – December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 6

Sildenafil Cream, 3.6% Concept Testing - Cream Formulation



It's not a pill. How do FSAD sufferers react to that?

- Pills tend to be synonymous with side effects, need to be taken regularly to be effective and their contents are metabolized by the liver because they enter the bloodstream.
- FSAD sufferers agree that these are the primary disadvantages of pills.





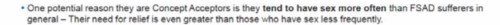
Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 10.

Sildenafil Cream, 3.6% Concept Testing - Concept Acceptors



What is the profile of *Concept Acceptors* (in terms of symptoms experienced, relationship to the condition of FSAD, etc.)?

- Concept Acceptors are not widely different from FSAD sufferers as a whole. However, they do appear to be the most severe sufferers of FSAD:
- The intensity of their symptoms and feelings is much stronger.
 - . They experience some of the physical symptoms more frequently, such as:
 - · Lack of genital wetness or lubrication;
 - * Lack of genital or clitoral fullness, pressure or engorgement;
 - · Lack of desire for intimacy and;
 - . Genital tenseness or tightness.
 - · Not surprisingly, they have an even deeper desire:
 - . For their bodies to respond to touch and intimacy, the way they used to;
 - * To have help becoming/staying physically aroused.
 - On an emotional level, they are even MORE....
 - . Bothered by their low sexual desire;
 - · Dissatisfied with their sex lives;
 - · Frustrated with their sexual problem;
 - . Engulfed by guilt about their sexual difficulties;
 - · Worried about sex;
 - . Likely to feel sexually inadequate/inferior and;
 - · Embarrassed.







Source data on File: Research Report – December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slide 11.

Sildenafil Cream, 3.6% Concept Testing – Purchase Interest

- 82% of respondents indicated they would be likely to purchase the product if it were currently available.
- A subgroup of respondents aged 35-54 had a higher purchase interest (86%) vs. the aggregate (82%).

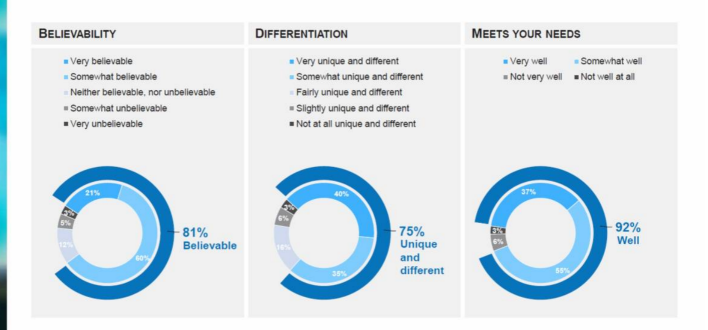
	Total n=402	Age group		FSAD LT SA 35-54	
		21-44 n=195	45-60 n=207	Yes n=120	No n=282
Likely	82%	81%	84%	86%	80%
Very likely	35%	31%	39%	46% 🛧	29% 4
Somewhatlikely	47%	50%	44%	40% 🔸	51% 1
Neither likely, nor unlikely	10%	9%	12%	10%	10%
Unlikely	8%	11% ↑	5% +	4%	10%
Somewhat unlikely	4%	6%	3%	2%	6%
Very unlikely	4%	5%	2%	2%	4%

* LT- in a long-term relationship SA - currently sexually active 35-54 - ages of 35 to 54



Source data on File: Research Report – December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slides 5-22

Sildenafil Cream, 3.6% Concept Testing – Believability & Viability





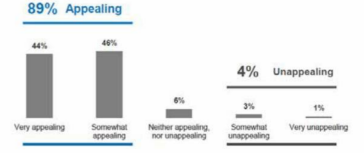
Source data on File: Research Report – December 2015; AD Hoc research - Sildenafii Topical Cream for Women Concept Test, slides 5-22.

Sildenafil Cream, 3.6% Concept Testing - Concept Appeal

- The majority of respondents (89%) considered the concept appealing.
- The largest proportion of respondents to consider the concept very appealing were women between the ages of 35-54.

Total n=402	Age group		FSAD LT SA 35-54*	
	21-44 n=195	45-60 n=207	Yes n=120	No n=282
89% 44%	88% 41%	91% 48%	95% + 50%	87% 4
46%	47%	44%	45%	46%
6%	6%	7%	4%	8%
4%	6% +	2% +	2%	6%
3%	4%	2%	196	4%
196	2% +	0% +	1%	2%
	n=402 89% 44% 46% 6% 4% 3%	21-44 n=195 89% 88% 44% 41% 46% 6% 6% 6% 4% 6% + 3% 4%	21-44 45-60 n=402 n=195 n=207	21-44 45-60 Yes n=120

LT- in a long-term relationship SA - currently sexually active 35-54 - ages of 35 to 54





Source data on File: Research Report - December 2015; AD Hoc research - Sildenafil Topical Cream for Women Concept Test, slides 5-22