Velva L. Price District Clerk Travis County D-1-GN-20-005184

Alexus Rodriguez

NO. D-1-GN-20-005184

THE STATE OF TEXAS,	§	IN THE DISTRICT COURT OF
	§	
Plaintiff,	§	
	§	
V.	§	TRAVIS COUNTY, TEXAS
	§	
C.R. BARD, INC.,	§	
	§	
Defendant.	8	200TH JUDICIAL DISTRICT

AGREED FINAL JUDGMENT AND PERMANENT INJUNCTION

Plaintiff, the State of Texas, has filed a petition in this action seeking a permanent injunction and other relief in this matter pursuant to the Texas Deceptive Trade Practices—Consumer Protection Act, Tex. Bus. & Com. Code Ann. §17.41 et seq. ("DTPA"), alleging that Defendant C. R. Bard, Inc. ("BARD" or "Defendant"), committed violations of the aforementioned Act. Plaintiff, by its counsel, and BARD, by its counsel, have agreed to the entry of this Agreed Final Judgment and Permanent Injunction ("Judgment") by the Court without trial or adjudication of any issue of fact or law, and without finding or admission of wrongdoing or liability of any kind.

IT IS HEREBY ORDERED THAT:

1. FINDINGS

- 1.1. This Court has jurisdiction over the subject matter of this lawsuit and over all Parties.
 - 1.2. The terms of this Judgment shall be governed by the laws of the State of Texas.
- 1.3. Entry of this Judgment is in the public interest and reflects a negotiated agreement among the Parties.

1.4. The Parties have agreed to resolve the issues resulting from the Covered Conduct

by entering into this Judgment.¹

1.5. BARD is willing to enter into this Judgment regarding the Covered Conduct in

order to resolve the Attorneys General's concerns under the State Consumer Protection Laws as

to the matters addressed in this Judgment and thereby avoid significant expense, inconvenience,

and uncertainty.

1.6. BARD is entering into this Judgment solely for the purpose of settlement, and

nothing contained herein may be taken as or construed to be an admission or concession of any

violation of law, rule, or regulation, or of any other matter of fact or law, or of any liability or

wrongdoing, all of which BARD expressly denies. BARD does not admit any violation of the

State Consumer Protection Laws set forth in footnote 4, and does not admit any wrongdoing that

was or could have been alleged by any Attorney General before the date of the Judgment under

those laws. No part of this Judgment, including its statements and commitments, shall constitute

evidence of any liability, fault, or wrongdoing by BARD.

1.7. This Judgment shall not be construed or used as a waiver or limitation of any

defense otherwise available to BARD in any other action, or of BARD's right to defend itself from,

or make any arguments in, any other private individual, regulatory, governmental, or class claims

or suits relating to the subject matter or terms of this Judgment. This Judgment is made without

trial or adjudication of any issue of fact or law or finding of liability of any kind. Notwithstanding

the foregoing, a State may file an action to enforce the terms of this Judgment.

¹This agreement is entered into pursuant to and subject to the State Consumer Protection laws cited in

footnote 4.

1.8. No part of this Judgment shall create a private cause of action or confer any right

to any third party for violation of any federal or state statute except that a State may file an action

to enforce the terms of this Judgment. It is the intent of the Parties that this Judgment shall not

be binding or admissible in any other matter, including, but not limited to, any investigation or

litigation, other than in connection with the enforcement of this Judgment.

1.9. This Judgment (or any portion thereof) shall in no way be construed to prohibit

BARD from making representations with respect to any BARD products in Labeling that are

required under Federal law, regulations, or policies or guidance having the force of law, including

in Food and Drug Administration ("FDA") approved Labeling.

1.10. Nothing in this Judgment shall require BARD to:

(a) take any action that is prohibited by the Food, Drug and Cosmetic Act, 21

U.S.C. § 301 et seq. ("FDCA") or any regulation promulgated thereunder,

or by the FDA; or

(b) fail to take any specific action that is expressly permitted or is required by

the FDCA or any regulation promulgated thereunder.

2. **DEFINITIONS**

The following definitions shall be used in construing the Judgment/Order:

2.1. "Covered Conduct" means BARD's marketing and promotional practices, and

dissemination of information to Health Care Providers (HCPs) and consumers, regarding

Urogynecologic Surgical Mesh products, including but not limited to the dissemination of

Marketing Materials, disclosure of Significant or Inherent Complications in Instructions for Use

(IFUs), Sponsorship of any programs, training any sales professionals, the publication of any

State of Texas v. C.R. Bard, Inc.
Agreed Final Judgment and Permanent Injunction

clinical or pre-clinical data, or the reporting of MDRs or adverse events, through the Effective

Date of the Judgment.

2.2. "Effective Date" means the date on which a copy of the Judgment, duly executed

by BARD and by the Signatory Attorney General, is approved by, and becomes a Judgment of the

Court.

2.3. "Health Care Provider" or "HCP" means any physician or other health care

practitioner who is licensed to provide health care services.

2.4. "BARD" means C. R. Bard, Inc. and Becton, Dickinson and Company and all of

their officers, directors, employees, representatives, agents, affiliates, parents, subsidiaries,

operating companies, assigns and successors.

2.5. "Labeling" means "all labels and other written, printed, or graphic matter (1) upon

any article or any of its containers or wrappers, or (2) accompanying such article," as defined under

Section 201(m) of the Federal Food, Drug, and Cosmetic Act (FDCA).

2.6. "Marketing Materials" means any written, electronic, or verbal material or

statements either publicly disseminated (including videos, websites it hosts or controls, or any

other form of media) or made for the purpose of public dissemination in the United States, in the

course of marketing, promoting, or informing Health Care Providers, consumers, or patients about

Urogynecologic Surgical Mesh, including, but not limited to, HCP training materials and training

materials for sales representatives made for the purpose of public dissemination and delivery to

HCPs.

2.7. "Multistate Executive Committee" means the Attorneys General and their staffs

representing California, Florida, Indiana, Maryland, Ohio, South Carolina, Texas, and

Washington.

2.8. "Multistate Working Group" means the Attorneys General and their staffs

representing Alabama, Alaska, Arizona, Arkansas, California, Colorado, Connecticut, Delaware,

District of Columbia, Florida, Georgia, Hawaii², Idaho, Illinois, Indiana, Iowa, Kansas, Kentucky,

Louisiana, Maine, Maryland, Massachusetts, Michigan, Minnesota, Mississippi, Missouri,

Montana, Nebraska, Nevada, New Hampshire, New Jersey, New Mexico, New York, North

Carolina, North Dakota, Ohio, Oklahoma, Oregon, Pennsylvania, Rhode Island, South Carolina,

South Dakota, Tennessee, Texas, Utah³, Vermont, Virginia, Washington, and Wisconsin.

2.9. "Parties" means BARD as defined in Section 2.4 and the Signatory Attorney

General.

2.10. "Post-effective Date Urogynecologic Surgical Mesh" means Urogynecologic

Surgical Mesh that enters the market in the United States after the Effective Date, and that is not

identical or substantially equivalent to Urogynecologic Surgical Mesh that was on the market in

the United States prior to the Effective Date.

2.11. "Significant Complications" means all complications of Urogynecologic Surgical

Mesh, including complications discovered subsequent to the Effective Date, which constitute

clinically significant risks material to a Health Care Provider's decision to implant Urogynecologic

Surgical Mesh.

²Hawaii is represented on this matter by its Office of Consumer Protection, an agency which is not part of the state Attorney General's Office, but which is statutorily authorized to undertake consumer protection functions, including legal representation of the State of Hawaii. For simplicity, the entire group will be referred to as the "Attorneys General," and such designation, as it includes Hawaii, refers to the Executive Director of the State of

Hawaii Office of Consumer Protection.

³With regard to Utah, the Utah Division of Consumer Protection is charged with administering and enforcing the Consumer Sales Practices Act, the statute relevant to this Judgment/Order. References to the "States," "Parties," or "Attorneys General," with respect to Utah, refers to the Utah Division of Consumer Protection.

- 2.12. "Inherent Mesh Complications" means Significant Complications that may not be eliminated with surgical technique and are associated with the use of Urogynecologic Surgical Mesh. Disclosure of such risks shall include an adequate description of the chronicity, acuteness, and permanence of the risks. A non-verbatim description of these risks shall include, but are not limited to, risks of:
 - Exposure of mesh material into the vagina, which can be associated with pain during intercourse for the woman and/or her partner
 - Pain caused by exposure may be severe and may result in permanent sexual dysfunction
 - Erosion
 - Implantation of Urogynecologic Surgical Mesh through the vagina may cause bacterial contamination
 - Infection
 - Voiding dysfunction, including de novo urge incontinence
 - Foreign body reaction
 - Inflammation
 - Scar plating around mesh
 - Clinical consequences of mesh contracture
 - Acute and/or chronic pain
 - Pelvic pain, which in some patients may not resolve
 - Pain with intercourse, which in some patients may not resolve
 - Excessive contraction or shrinkage of the tissue surrounding the mesh, vaginal scarring, tightening and/or shortening may occur

Such description shall also note that the occurrence of one or more of these complications may require treatment or surgical intervention:

- In some instances, the complication may persist as a permanent condition after the surgical intervention or other treatment;
- Removal of mesh or correction of mesh-related complications may involve multiple surgeries; and
- iii. Complete removal of mesh may not be possible and additional surgeries may not always fully correct the complications

However, for Post-Effective Date Urogynecologic Surgical Mesh, a non-verbatim description of these risks may include, but are not limited to, the risks listed in the bullet points above, depending upon the available Valid Scientific Evidence.

- 2.13. "Signatory Attorney General" means the Attorney General of Texas, or his authorized designee, who has agreed to this Judgment.
- 2.14. "Sponsor" or "Sponsorship" means to pay for in whole or in part, to provide financial support or subsidization, or to provide goods or materials of value in support, but does not include de minimis support.
- 2.15. "State Consumer Protection Laws" means the consumer protection laws cited in footnote 4 under which the Attorneys General have conducted the investigation.⁴

⁴ALABAMA – Alabama Deceptive Trade Practices Act § 8-19-1 et seq. (2002); ALASKA – Alaska Unfair Trade Practices and Consumer Protection Act AS 45.50.471 – 45.50.561; ARIZONA - Consumer Fraud Act, A.R.S. §44-1521 et seq.; ARKANSAS - Arkansas Deceptive Trade Practices Act, Ark. Code Ann. § 4-88-101, et seq.; CALIFORNIA - Bus. & Prof Code §§ 17200 et seq. and 17500 et seq.; COLORADO - Colorado Consumer Protection Act, Colo. Rev. Stat. § 6-1-101 et seq.; CONNECTICUT - Connecticut Unfair Trade Practices Act, Conn. Gen Stat. §§ 42-110a through 42-110q; DELAWARE – Delaware Consumer Fraud Act, Del. CODE ANN. tit. 6, §§ 2511 to 2527; DISTRICT OF COLUMBIA, District of Columbia Consumer Protection Procedures Act, D.C. Code §§ 28-3901 et seq.; FLORIDA - Florida Deceptive and Unfair Trade Practices Act, Part II, Chapter 501, Florida Statutes, 501.201 et. seq.; GEORGIA - Fair Business Practices Act, O.C.G.A. Sections 10-1-390 et seq.; HAWAII - Uniform Deceptive Trade Practice Act, Haw. Rev. Stat. Chpt. 481A and Haw. Rev. Stat. Chpt. 480; IDAHO - Idaho Consumer Protection Act, Idaho Code § 48-601 et seq.; ILLINOIS – Consumer Fraud and Deceptive Business Practices Act, 815 ILCS 505/2 et seq.; INDIANA - Deceptive Consumer Sales Act, Ind. Code §§ 24-5-0.5-0.1 to 24-5-0.5-12; IOWA -Iowa Consumer Fraud Act, Iowa Code Section 714.16; KANSAS - Kansas Consumer Protection Act, K.S.A. 50-623 et seq.; KENTUCKY - Kentucky Consumer Protection Act, KRS Ch. 367.110, et seq.; LOUISIANA - Unfair Trade-Practices and Consumer Protection Law, LSA-R.S. 51:1401, et seq.; MAINE - Unfair Trade Practices Act, 5 M.R.S.A. § 207 et seg.; MARYLAND - Maryland Consumer Protection Act, Md. Code Ann., Com. Law §§ 13-101 et seg.; MASSACHUSETTS - Mass. Gen. Laws c. 93A, §§ 2 and 4; MICHIGAN - Michigan Consumer Protection Act, MCL § 445.901 et seq.; MINNESOTA – Minn. Stat. §§325D.44, 325F.69; MISSISSIPPI - Mississippi Consumer Protection Act, Miss. Code Ann. § 75-24-1, et seq.; MISSOURI – Missouri Merchandising Practices Act, Mo. Rev. Stat. §§ 407.010 et seq.; MONTANA – Montana Consumer Protection Act §§ 30-14-101 et seq.; NEBRASKA – Consumer Protection Act, Neb. Rev. Stat. §§ 59-1601 et seq. and Uniform Deceptive Trade Practices Act, Neb. Rev. Stat. §§ 87-301 et seq.; NEVADA - Deceptive Trade Practices Act, Nevada Revised Statutes 598.0903 et seq.; NEW HAMPSHIRE - NH RSA §358-A et seq; NEW JERSEY - New Jersey Consumer Fraud Act, NJSA 56:8-1 et seq.; NEW MEXICO - NMSA 1978, § 57-12-1 et seg.; NEW YORK - General Business Law Art. 22-A, §§ 349-50, and Executive Law § 63(12); NORTH CAROLINA – North Carolina Unfair and Deceptive Trade Practices Act, N.C.G.S. 75-1.1, et seq.; NORTH DAKOTA - Unlawful Sales or Advertising Practices, N.D. Cent. Code § 51-15-02 et seq.; OHIO - Ohio Consumer Sales Practices Act, R.C. 1345.01, et seq.; OKLAHOMA - Oklahoma Consumer Protection Act 15 O.S. §§ 751 et seq.; OREGON – Oregon Unlawful Trade Practices Act, Or. Rev. Stat. § 646.605 et seq.; PENNSYLVANIA - Pennsylvania Unfair Trade Practices and Consumer Protection Law, 73 P.S. 201-1 et seq.;

2.16. "Urogynecologic Surgical Mesh" means any medical device cleared or approved

by the FDA (as the term "device" is defined in 21 U.S.C. § 321(h)) that contains synthetic, multi-

strand, knitted, or woven mesh and that is indicated to be used for implantation in the pelvic floor

to treat stress urinary incontinence (SUI) and/or pelvic organ prolapse (POP) sold or marketed in

the United States.

2.17. "Valid Scientific Evidence" means evidence from well-controlled investigations,

partially controlled studies, studies and objective trials without matched controls, well-

documented case histories conducted by qualified experts, or reports of significant human

experience with a marketed device, from which it can fairly and responsibly be concluded by

qualified experts that there is reasonable assurance to substantiate that a representation is true.

2.18. Any reference to a written document shall mean a physical paper copy of the

document, electronic version of the document, or electronic access to such document.

3. COMPLIANCE PROVISIONS

A. Exit from Urogynecologic Surgical Mesh Business

3.1. BARD states that it ceased the marketing, promotion, sale, and distribution of

Urogynecologic Surgical Mesh in the United States and the manufacturing of Urogynecologic

Surgical Mesh for sale in the United States by December 30, 2016.

RHODE ISLAND – Deceptive Trade Practices Act, Rhode Island Gen. Laws § 6-13.1-1, et seq.; SOUTH CAROLINA – South Carolina Unfair Trade Practices Act, S.C. Code Ann. § 39-5-10 et seq.; SOUTH DAKOTA – South Dakota Deceptive Trade Practices and Consumer Protection, SDCL ch. 37-24; TENNESSEE – Tennessee Consumer Protection Act, Tenn. Code Ann. 47-18-101 et seq.; TEXAS – Texas Deceptive Trade Practices-Consumer Protection Act, Tex. Bus. And Com. Code 17.41, et seq.; UTAH - Consumer Sales Practices Act, Utah Code Ann. §§ 13-11-1 et seq.; VERMONT – Vermont Consumer Protection Act, 9 V.S.A. § 2451, et seq.; VIRGINIA-Virginia Consumer Protection Act, Va Code Ann. §59.1-196 et seq.; WASHINGTON – Unfair Business Practices/Consumer Protection

Act, RCW §§ 19.86 et seq.; WISCONSIN – Wis. Stat. § 100.18 (Fraudulent Representations).

3.2. In the event that BARD engages in any conduct involving the manufacture,

promotion, marketing, sale, or distribution of Urogynecologic Surgical Mesh, either directly or

indirectly through any third parties, in the United States, it shall be bound by the following

provisions contained in Sections 3.4 through 3.27 of this Judgment/Order for ten (10) years from

the date of first sale of an Urogynecologic Surgical Mesh product in the United States or for twenty

(20) years from the Effective Date of this Agreement, whichever is less. Section 3.3 is not time

restricted. Nothing in this Judgment/Order shall be construed to require BARD, for any

Urogynecologic Surgical Mesh product approved through the FDA Premarket Approval process,

to utilize product labeling different from that which is approved by the FDA.

B. Marketing, Information, and Training

3.3. In promoting Urogynecologic Surgical Mesh, BARD shall not violate the Texas

Deceptive Trade Practices-Consumer Protection Act, TEX. BUS. & COM. CODE §§17.41 et seq.

3.4. BARD shall not, in any Marketing Materials, make any claim comparing safety or

efficacy clinical outcomes with the use of Urogynecologic Surgical Mesh to any non-mesh

procedure safety or efficacy clinical outcomes, unless any such representation is supported by

Valid Scientific Evidence. BARD, however, may make comparisons in any Marketing Materials

not involving safety or efficacy clinical outcomes, if not false, misleading, or deceptive.

3.5. BARD shall not, in any Marketing Materials, misrepresent the safety or efficacy of

its Urogynecologic Surgical Mesh by omitting Significant Complications or Inherent Mesh

Complications, as appropriate given the length, context, medium, and placement of the Marketing

Material and in all instances where the Marketing Material purports to address the subject of

complications.

3.6. In any Marketing Material that is intended to reach patients or consumers other than

or in addition to Health Care Providers, BARD shall also include descriptions of Significant

Complications and Inherent Mesh Complications in terms reasonably understandable to a patient.

3.7. BARD shall not, in any Marketing Materials, misrepresent the extent to which

Inherent Mesh Complications are risks or complications common to all pelvic floor or other

surgeries.

3.8. BARD shall not, in any Marketing Materials, represent or imply that Significant

Complications or Inherent Mesh Complications can be eliminated with surgical experience or

technique alone. However, for Post-Effective Date Urogynecologic Surgical Mesh, BARD may,

in any Marketing Materials, represent or imply that Significant Complications can be eliminated

with surgical experience or technique alone, if such statement is supported by Valid Scientific

Evidence.

3.9. BARD shall not represent or imply that such Urogynecologic Surgical Mesh does

not cause a foreign body reaction, including any chronic foreign body reaction, after the

Urogynecologic Surgical Mesh is implanted inside the body. However, for Post-Effective Date

Urogynecologic Surgical Mesh, BARD may represent or imply that such Urogynecologic Surgical

Mesh does not cause a foreign body reaction, including any chronic foreign body reaction, after

the Urogynecologic Surgical Mesh is implanted inside the body, if such statement is supported by

Valid Scientific Evidence.

3.10. BARD shall not, in any Marketing Materials, represent or imply that such

Urogynecologic Surgical Mesh is "soft" or that it has "multidirectional elasticity" within the body

after implantation or use any other phrases having an equivalent meaning. However, for Post-

Effective Date Urogynecologic Surgical Mesh, BARD may, in any Marketing Materials, represent

or imply that such Urogynecologic Surgical Mesh is "soft" or that it has "multidirectional

elasticity" within the body after implantation or use any other phrases having an equivalent

meaning, if such statement is supported by Valid Scientific Evidence. Nothing shall prevent

BARD from making claims to Health Care Providers about the softness and elasticity of

Urogynecologic Surgical Mesh prior to implantation inside the body provided the claims do not

suggest these properties are retained in the body.

3.11. BARD shall not, in any Marketing Materials, represent or imply that such

Urogynecologic Surgical Mesh, including its collagen Urogynecologic Surgical Mesh, helps the

body more readily accept a foreign body implant, or reduces the risk of foreign body reaction,

erosion, infection, or any other Urogynecologic Surgical Mesh complications, including any

Significant Complications or Inherent Complications. However, for Post-Effective Date

Urogynecologic Surgical Mesh, BARD may, in any Marketing Materials, represent or imply that

such Urogynecologic Surgical Mesh, including its collagen Urogynecologic Surgical Mesh, helps

the body more readily accept a foreign body implant, or reduces the risk of foreign body reaction,

erosion, infection, or any other Urogynecologic Surgical Mesh complications, including any

Significant Complications or Inherent Complications, if such statement is supported by Valid

Scientific Evidence.

3.12. BARD shall not, in any Marketing Materials, misrepresent the FDA approval or

clearance status of its Urogynecologic Surgical Mesh devices or the extent to which any of its

Urogynecologic Surgical Mesh products have been studied or clinically proven.

3.13. BARD shall not, in any Marketing Materials, misrepresent the complexity of

Urogynecologic Surgical Mesh implantation procedures or the level of surgical skill and/or

experience necessary to perform these procedures safely. Moreover, BARD employees shall not

encourage a Health Care Provider to perform Urogynecologic Surgical Mesh implants without

receiving adequate information and training on how to implant its Urogynecologic Surgical Mesh.

3.14. In any training in which BARD provides risk information, either directly or through

third parties, to any Health Care Provider, BARD shall disclose all Significant Complications and

Inherent Mesh Complications of its Urogynecologic Surgical Mesh.

3.15. BARD shall, in the marketing and promotion of any Urogynecologic Surgical Mesh

product, ensure that its Marketing Materials and other communications do not misrepresent FDA

updates or communications regarding Urogynecologic Surgical Mesh.

C. Disclosures to Health Care Providers

3.16. To the extent not prohibited by federal law, BARD shall ensure that all IFUs for its

Urogynecologic Surgical Mesh products cleared through the 510(k) process include a list of all

known Significant Complications and Inherent Mesh Complications.

3.17. BARD shall evaluate emerging risk information on an ongoing basis and, consistent

with such risk information, shall update the warnings and precautions section of IFUs and all

Marketing Material to include Significant Complications associated with its Urogynecologic

Surgical Mesh products as soon as practicable. If Bard obtains, receives, or is aware of any new

risk information that necessitates a more immediate disclosure for public health and safety

purposes, Bard shall notify HCPs of this information through other means, such as notices or "dear

doctor letters," as appropriate given the nature of the new information and unless otherwise

directed by the FDA.

State of Texas v. C.R. Bard, Inc.
Agreed Final Judgment and Permanent Injunction

D. Studies, Clinical Data, and Sponsorship

3.18. BARD shall, when citing to any clinical study, clinical data, or preclinical data,

present a fair and balanced view of available scientific literature with respect to the safety, efficacy,

risks and complications of Urogynecologic Surgical Mesh.

3.19. BARD shall, when citing to any clinical study, clinical data, or preclinical data

regarding Urogynecologic Surgical Mesh in its Marketing Materials, not misrepresent the results,

scope, or clinical significance of any particular clinical study, clinical data, or preclinical data,

including by implying a more favorable result than supported by the study or data.

3.20. BARD shall, when submitting a clinical study, clinical data, or preclinical data

regarding Urogynecologic Surgical Mesh for publication, disclose BARD's role as a Sponsor and

any author's potential conflict of interest consistent with the disclosure requirements for the

International Committee of Medical Journal Editors (ICMJE) or, if different, the disclosure

policies of the relevant publication.

3.21. BARD shall not cite to any clinical study, clinical data, or preclinical data regarding

Urogynecologic Surgical Mesh for which BARD has not complied with the requirements of

Section 3.D.

3.22. BARD shall not cite to any clinical study, clinical data, or preclinical data regarding

Urogynecologic Surgical Mesh for which any author/consultant, to the extent BARD knows, has

not complied with the applicable publication's conflict disclosure requirements unless BARD

discloses the conflict in a clear and conspicuous manner when citing to such study or data.

3.23. In all contracts for consulting services regarding Urogynecologic Surgical Mesh

between BARD and any Health Care Provider or other author/consultant, BARD shall include a

Sponsorship disclosure provision under which the Health Care Provider or other author/consultant

agrees that he or she shall, in terms likely to be read and understood by the audience, disclose in

any public presentation or submission for publication BARD's sponsorship of the contracted-for

activities. BARD shall also include a disclosure clause under which the Health Care Provider or

other author/consultant acknowledges that BARD may publicly report the fact that BARD made

value transfers to him or her. To the extent within its control, BARD shall ensure that any HCP

or author/consultant who submits for publication a clinical study, clinical data, or pre-clinical data

that BARD has Sponsored, authored, or edited, in whole or in part, shall comply with the

publication's conflict disclosure requirements.

3.24. In accordance with applicable law, BARD shall register BARD-sponsored clinical

studies regarding its Urogynecologic Surgical Mesh with ClinicalTrials.gov. BARD shall also

retain any design history files and clinical records, including but not limited to clinical data,

relating to its post-December 30, 2016 Urogynecologic Surgical Mesh devices and any

Urogynecologic Surgical Mesh devices that existed prior to December 30, 2016 (or substantially

equivalent to such devices) over which it has or should have possession, custody or control for 15

years past the last sale date of the Urogynecologic Surgical Mesh devices to which those files and

records apply, unless a longer period is required by applicable law. BARD shall retain any non-

clinical data relating to its post-December 30, 2016 Urogynecologic Surgical Mesh devices and

any Urogynecologic Surgical Mesh devices that existed prior to December 30, 2016 (or

substantially equivalent to such devices) over which it has or should have possession, custody or

control until December 30, 2031, if not introduced prior to that date. If introduced prior to

December 30, 2031, then BARD shall retain non-clinical data for 15 years past the last sale date,

unless a longer period is required by applicable law.

State of Texas v. C.R. Bard, Inc.

Agreed Final Judgment and Permanent Injunction

Page 14 of 24

E. BARD Internal Policies and Training

3.25. BARD shall ensure that its independent contractors, agents, and employees, who sell, market, or promote Urogynecologic Surgical Mesh or otherwise train, provide information to, or communicate with Health Care Providers regarding Urogynecologic Surgical Mesh, are adequately informed and trained regarding their obligations to report all patient complaints and/or

adverse events to BARD.

3.26. BARD shall ensure that its company practices regarding the reporting of patient complaints relating to Urogynecologic Surgical Mesh as MDR reportable adverse events are consistent with FDA requirements.

F. Monitoring and Compliance

3.27. BARD shall be responsible for ensuring monitoring and compliance with the provisions of this Judgment.

4. PAYMENT

4.1. BARD shall pay a total amount of \$60 million as follows: 1) the initial payment of \$15 million shall be paid by the later of October 30, 2020 or 30 days after the Effective Date; 2) the second payment of \$15 million shall be paid by April 1, 2021; and 3) the final payment of \$30 million shall be paid by October 30, 2021. These payments will be divided and paid by BARD to each Signatory Attorney General of the Multistate Working Group in amounts to be designated by and in the sole discretion of the Multistate Executive Committee. Said payments shall be used

⁵ Bard shall pay Texas \$887,719.75 by the later of October 30, 2020 or 30 days after the Effective Date; \$887,719.75 by April 1, 2021; and \$1,775,499.50 by October 30, 2021. Of these amounts, \$2,485,615.30 will be allocated as a payment to the State of Texas pursuant to Texas Government Code Section 402.007, and \$1,065,263.70 will be allocated as attorneys' fees and investigative costs under Texas Government Code Section 402.006(c).

by the States as attorneys' fees and other costs of investigation and litigation, or to be placed in,

or applied to, the consumer protection enforcement including future consumer protection

enforcement, consumer education, litigation or local consumer aid fund or revolving fund, used to

defray the costs of the inquiry leading hereto, or for any lawful purpose, at the sole discretion of

each Signatory Attorney General. The Parties acknowledge that the payments described herein

are not a fine, penalty, or payment in lieu thereof.

5. ENFORCEMENT

5.1. For the purposes of resolving disputes with respect to compliance with this

Judgment, should any of the Signatory Attorneys General have a reasonable basis to believe that

BARD has engaged in a practice that violates a provision of this Judgment subsequent to the

Effective Date, then such Attorney General shall notify BARD in writing of the specific objection,

identify with particularity the provision of this Judgment that the practice appears to violate, and

give BARD thirty (30) days to respond to the notification; provided, however, that a Signatory

Attorney General may take any action if the Signatory Attorney General believes that, because of

the specific practice, a threat to the health or safety of the public requires immediate action.

5.2. Upon receipt of written notice, BARD shall provide a good-faith written response

to the Attorney General notification, containing either a statement explaining why BARD believes

it is in compliance with the Judgment, or a detailed explanation of how the alleged violation

occurred and a statement explaining how BARD intends to remedy the alleged breach. Nothing

in this section shall be interpreted to limit the State of Texas's Civil Investigative Demand ("CID")

or investigative subpoena authority, to the extent such authority exists under applicable law, and

BARD reserves all of its rights in responding to a CID or investigative subpoena issued pursuant

to such authority.

5.3. The Attorney General may agree, in writing, to provide BARD with additional time

beyond the thirty (30) days to respond to a notice provided under section 5.1 above.

5.4. Upon giving BARD thirty (30) days to respond to the notification described above,

the Signatory Attorney General shall also be permitted reasonable access to inspect and copy

relevant, non-privileged, non-work product records and documents in the possession, custody, or

control of BARD that relate to BARD's compliance with each provision of this Judgment pursuant

to that State's CID or investigative subpoena authority. If the Signatory Attorney General makes

or requests copies of any documents during the course of that inspection, the Signatory Attorney

General shall provide a list of those documents to BARD.

5.5. The State may assert any claim that BARD has violated this Judgment in a separate

civil action to enforce compliance with this Judgment, or may seek any other relief afforded by

law for violations of the Judgment, but only after providing BARD an opportunity to respond to

the notification described in paragraph 5.1 above; provided, however, that a Signatory Attorney

General may take any action if the Signatory Attorney General believes that, because of the

specific practice, a threat to the health or safety of the public requires immediate action.

6. RELEASE

6.1. Released Claims. By its execution of this Judgment, the State of Texas releases

and forever discharges BARD and its past and present officers, directors, employees,

representatives, agents, affiliates, parents, subsidiaries, operating companies, predecessors, assigns

and successors (collectively, the "Releasees") from the following: all civil causes of action, claims,

damages, restitution, disgorgement, fines, costs, attorney's fees, or penalties that the Texas

Attorney General has asserted or could have asserted against the Releasees under the State

Consumer Protection Laws, or any amendments thereto, or by common law claims concerning

deceptive or fraudulent trade practices, that the Signatory Attorney General has the authority to release resulting from the Covered Conduct up to and including the Effective Date. For purposes of this Section 6.1, Releasees do not include Covidien Ltd. or Medtronic PLC, or their past and present officers, directors, employees, representatives, agents, affiliates, parents, subsidiaries, operating companies, predecessors, assigns and successors.

- 6.2. <u>Claims Not Covered.</u> Notwithstanding any term of this Judgment, specifically reserved and excluded from the release in Paragraph 6.1 as to any entity or person, including Releasees, are any and all of the following:
 - (a) Any criminal liability that any person or entity, including Releasees, has or may have to the State of Texas;
 - (b) Any civil or administrative liability that any person and/or entity, including Releasees, has or may have to the State of Texas not expressly covered by the release in Section 6.1, including, but not limited to, any and all of the following claims:
 - i. State or federal antitrust violations;
 - ii. Claims involving "best price," "average wholesale price,""wholesale acquisition cost," or any reporting practices;
 - iii. Medicaid claims, including but not limited to federal Medicaid drug rebate statute violations, Medicaid fraud or abuse (whether common law, statutory or otherwise), and/or kickback violations related to any state's Medicaid program;
 - iv. State false claims violations; and
 - v. Claims to enforce the terms and conditions of this Judgment.

- (c) Actions of, or on behalf of, state program payors of the State of Texas arising from the purchase of Urogynecologic Surgical Mesh.
- (d) Any claims individual consumers have or may have under above-cited

 State Consumer Protection Laws against any person or entity, including the Releasees.
- 6.3. Nothing contained in this Judgment shall relieve BARD of the obligations it maintains under any other Judgment or agreement relating to any BARD product.

7. ADDITIONAL PROVISIONS

- 7.1. Nothing in this Judgment shall be construed to authorize or require any action by BARD in violation of applicable federal, state, or other laws.
- 7.2. <u>Modification.</u> The Judgment may be modified by a stipulation of the Parties as approved by the Court, or by court proceedings resulting in a modified judgment of the Court.
- 7.3. BARD shall not cause or encourage third parties, nor knowingly permit third parties acting on its behalf, to engage in practices from which BARD is prohibited by this Judgment.
- 7.4. The acceptance of this Judgment by the State of Texas shall not be deemed approval by the State of Texas of any of BARD's advertising or business practices. Further, neither BARD nor anyone acting on its behalf shall state or imply, or cause to be stated or implied, that the State of Texas or any other governmental unit of Texas has approved, sanctioned or authorized any practice, act, advertisement, or conduct of BARD.
- 7.5. Any failure by any party to this Judgment to insist upon the strict performance by any other party of any of the provisions of this Judgment shall not be deemed a waiver of any of the provisions of this Judgment, and such party, notwithstanding such failure, shall have the right thereafter to insist upon the specific performance of any and all of the provisions of this Judgment.

7.6. Entire Agreement: This Judgment represents the full and complete terms of the

settlement entered into by the Parties hereto. In any action undertaken by the Parties, no prior

versions of this Judgment and no prior versions of any of its terms that were not entered by the

Court in this Judgment, may be introduced for any purpose whatsoever.

7.7. Jurisdiction: This Court retains jurisdiction of this Judgment and the Parties hereto

for the purpose of enforcing and modifying this Judgment and for the purpose of granting such

additional relief as may be necessary and appropriate.

7.8. Counterparts: This Judgment may be executed in counterparts, and a facsimile or

.pdf signature shall be deemed to be, and shall have the same force and effect as, an original

signature.

7.9. Notice: All Notices under this Judgment shall be provided to the following via

email and Overnight Mail:

Defendant:

Greg A. Dadika

Senior Vice President, Chief Legal Counsel

Becton Dickinson and Company

Greg.Dadika@bd.com

Copy to BARD's attorneys at

Troutman Pepper via electronic mail sent to:

Barry H. Boise (boiseb@troutman.com)

For the State:

Nanette DiNunzio

Assistant Attorney General

Consumer Protection Division

P. O. Box 12548

Austin, TX 78711-2548

Nanette.DiNunzio@oag.texas.gov

7.10. To the extent that any provision of this Judgment obligates BARD to change any policy(ies) or procedure(s) and to the extent not already accomplished, BARD shall implement the

 $policy (ies) \ or \ procedure (s) \ as \ soon \ as \ reasonably \ practicable, \ but \ no \ later \ than \ 120 \ days \ after \ the$

Effective Date of this Judgment.

Dustin M. Howell

SO ORDERED, ADJUDGED AND DECREED

	September 29, 20.
PRESIDING JUDGE	Date

AGREED AS TO FORM AND SUBSTANCE AND ENTRY REQUESTED:

For Plaintiff State of Texas

Dated: Sept. 17, 2020

KEN PAXTON

Attorney General of Texas

JEFFREY C. MATEER

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RYAN L. BANGERT

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ATTORNEYS FOR THE STATE OF TEXAS

AGREED AS TO FORM AND SUBSTANCE AND ENTRY REQUESTED:

For Defendant C. R. Bard

GREG A. DADIKA

Senior Vice President, Chief Legal Counsel

Becton Dickinson and Company

Greg.Dadika@bd.com

AGREED AS TO FORM AND SUBSTANCE AND ENTRY REQUESTED:

Dated: September 22, 2020

VIRGINIA BELL FLYNN

Texas State Bar No. 24101258

Troutman Pepper Hamilton Sanders LLP

Virginia Ber Fran

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ATTORNEYS FOR DEFENDANT

Automated Certificate of eService

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Status as of 9/30/2020 2:08 PM CST

Case Contacts

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