

**IN THE UNITED STATES DISTRICT COURT
FOR THE SOUTHERN DISTRICT OF WEST VIRGINIA**

CHARLESTON DIVISION

IN RE: COLOPLAST CORP. PELVIC SUPPORT
SYSTEMS PRODUCTS LIABILITY LITIGATION

MDL 2387

THIS DOCUMENT RELATES TO
ALL CASES INCLUDING COLOPLAST WAVE 1 CASES

PRETRIAL ORDER # 105

(Order re: Amending PTO # 12 and
Adopting Plaintiff Fact Sheets and Defendant Fact Sheets)

PTO # 12 (Plaintiff Profile Forms, Plaintiff Fact Sheet, and Defendant Fact Sheets) contains provisions related to the Plaintiff Fact Sheet (“PFS”) and the Defendant Fact Sheet (“DFS”). It is **ORDERED** that PTO # 12 is amended to omit paragraphs 2.a, 3.a (first paragraph only) and 3.b. In light of the court’s recent decision to enter wave orders in this MDL, those provisions referring to bellwether cases and subgroups of bellwether cases are no longer applicable. However, the remaining provisions in PTO # 12 related to Plaintiff Profile Forms, PFSs and DFSs remain in place.

The parties have agreed to and submitted for entry, a proposed PFS with Verifications and Authorizations¹ and a proposed DFS, attached as Exhibits A and B respectively. The court adopts the PFS and DFS for use in the Coloplast MDL wave and other cases, where applicable, and it is **ORDERED** as follows:

¹ The PFS, Verifications and Authorizations can be found on the court’s website at www.wvsd.uscourts.gov under the Coloplast MDL, Plaintiff Fact Sheet. The Authorizations are the same as those used for the Plaintiff Profile Form.

(1) The PFSs, Verification(s) and Authorizations must be submitted to the plaintiffs' leadership and Coloplast and/or Mentor counsel (where applicable) at the following addresses electronically:

Plaintiffs - rburnett@rburnettlaw.com - (Riley Burnett)
lcausey@salim-beasley.com – (Lisa Causey-Streete)
jmostyn@mostynlaw.com – (Steve Mostyn)
mholbrook@holbrooklaw.com – (Mark Holbrook); and
crcallahan@mostynlaw.com - (Cossette Callahan)

Coloplast - ColoplastMesh@fulbright.com - (Lana Varney)

Mentor - MentorPPFs@tuckerellis.com - (Dustin Rawlin)

(2) Defendant(s) must submit a completed DFS where required to the plaintiffs electronically at:

rburnett@rburnettlaw.com - (Riley Burnett)
lcausey@salim-beasley.com – (Lisa Causey-Streete)
jmostyn@mostynlaw.com – (Steve Mostyn)
mholbrook@holbrooklaw.com – (Mark Holbrook); and
crcallahan@mostynlaw.com - (Cossette Callahan)

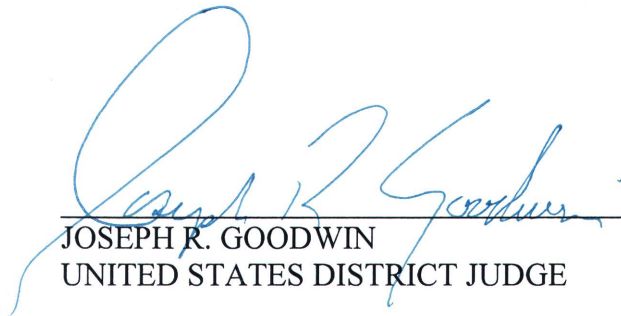
and defendant(s) must serve a completed DFS in each individual case where required on the individual plaintiff's counsel in that particular case.

(3) Any party who fails to comply with the PFS and DFS obligations under this order must comply with the provisions contained in PTO # 12 at paragraphs 2.c and 3.d.

The court **DIRECTS** the Clerk to file a copy of this order in 2:12-md-2387 and in the Coloplast Wave 1 Cases. In cases subsequently filed in this district after 2:16-cv-05443, a copy of the most recent pretrial order will be provided by the Clerk to counsel appearing in each new action at the time of filing of the complaint. In cases subsequently removed or transferred to this Court, a copy of the most recent pretrial order will be provided by the Clerk to counsel appearing in each new action upon removal or transfer. It shall be the responsibility of the parties to review and abide

by all pretrial orders previously entered by the Court. The orders may be accessed through the CM/ECF system or the Court's website at www.wvsd.uscourts.gov.

ENTER: June 21, 2016



JOSEPH R. GOODWIN
UNITED STATES DISTRICT JUDGE

**IN THE UNITED STATES DISTRICT COURT
FOR THE SOUTHERN DISTRICT OF WEST VIRGINIA**

CHARLESTON DIVISION

IN RE: COLOPLAST CORP.
PELVIC SUPPORT SYSTEMS
PRODUCTS LIABILITY
LITIGATION

MDL No. 2387

THIS DOCUMENT RELATES TO

Civil Action No.: _____

Name of Plaintiff

PLAINTIFF FACT SHEET

Each plaintiff who allegedly suffered injury as a result of a pelvic mesh product manufactured or sold by Coloplast Corp. must complete this Plaintiff Fact Sheet. In completing this Fact Sheet, you are under oath and must answer every question and provide information that is true and correct to the best of your knowledge. If you cannot recall all of the details requested, please provide as much information as you can and then state that your answer is incomplete and explain why as appropriate. If you select an "I Don't Know" answer, please state all that you do know about that subject. If any information you need to complete any part of the Fact Sheet is in the possession of your attorney, please consult with your attorney so that you can fully and accurately respond to the questions set out below. If you are completing the Fact Sheet for someone who cannot complete the Fact sheet herself, please answer as completely as you can.

The Fact Sheet shall be completed in accordance with the requirements and guidelines set forth in the applicable Case Management Order. A completed Fact Sheet shall be considered interrogatory answers pursuant to Fed. R. Civ. P. 33 and 34 and will be governed by the standards applicable to written discovery under Fed. R. Civ. P. 26 through 37. You must supplement your responses if you learn that they are incomplete or incorrect in any material respect. The questions and requests for production contained in the Fact Sheet are non-objectionable and shall be answered without objection. This Fact Sheet shall not preclude Defendants from seeking additional documents and information on a reasonable, case-by-case basis pursuant to the Federal Rules of Civil Procedure and as permitted by the applicable Case Management Order.

In filling out this form, please use the following definition: "healthcare provider" means any doctor, physician, surgeon, pharmacist, hospital, clinic, center, physician's office, infirmary, medical or diagnostic laboratory, or other facility that provides medical care or advice, and any pharmacy, x-ray department, radiology department, laboratory, physical therapist or physical therapy department, rehabilitation specialist, chiropractor, or other persons or entities involved in the diagnosis, care and/or treatment of you.

In filling out this form, the terms "You" or "Your" refer to the person who received pelvic mesh product(s) manufactured or sold by Coloplast Corp. and who is identified in Question I.1 (a) below.

To the extent that the form does not provide enough space to complete your responses or answers, please attach additional sheets as necessary.

I. BACKGROUND INFORMATION

- 1) Please state:
 - a. Full name of the person who received the pelvic mesh product(s), including maiden name:

 - b. Full name of the person completing this form, if different from the person listed in 1 (a) above, and the relationship of the person completing this form to the person listed in 1 (a) above: _____
 - c. The name and address of your primary attorney: _____
- 2) Your Social Security Number: _____
- 3) Your date of birth: _____
- 4) Your current residence address: _____

If you have lived at this address for less than 10 years, provide each of your prior residence addresses from 2000 to the present:

Prior Address	Dates You Lived At This Address

- 5) Have you ever been married? **Yes** ____ **No** ____

If yes, provide the names and addresses of each spouse and the inclusive dates of your marriage to each person.

- 6) Do you have children? **Yes** ____ **No** ____

If Yes, please provide the following information with respect to each child:

Full Name of Child	Date of Birth	Home Address (if different from yours)	Whether Biological/Adopted

- 7) Identify the name and age of any person who currently resides with you and their relationship to you:

- 8) Identify all secondary and post-secondary schools you attended, starting with high school and please provide the following information with respect to each:

Name of School	Address	Dates of Attendance	Degree Awarded	Major or Primary Field

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- 9) Please provide the following information for your employment history over the past 10 years up until the present:

Employer Name	Addresses	Job Title/ Description of Duties	Dates of Employment	Salary/Rate of Pay

- 10) Have you ever served in any branch of the military? **Yes** ____ **No** ____

If Yes, please provide the following information:

- a. Branch and dates of service, rank upon discharge and the type of discharge you received: _____

- b. Were you discharged from the military at any time for any reason relating to your medical, physical, or psychiatric condition? **Yes** ____ **No** ____

If Yes, state what that condition was: _____

- 11) Within the last ten years, have you been convicted of, or plead guilty to, a felony and/or crime of fraud or dishonesty? **Yes** ____ **No** ____

If Yes, please set forth where, when and the felony and/or crime:

II. CLAIM INFORMATION

- 1) Please complete the following chart for each implanted Coloplast Corp. pelvic mesh product. Insert additional lines as necessary.

Pelvic Mesh Product <u>and</u> lot number (if sticker affixed, so indicate)	Date of Implant and Name and Location of Implanting Facility	Reason for Implant	Implanting Doctor and Address
<u>Product No. 1:</u>			
<u>Product No. 2:</u>			
<u>Product No. 3:</u>			

- 2) For each pelvic mesh product identified above, describe your understanding of the medical condition for which you received the pelvic mesh product(s), including a detailed description of the symptoms, if any, for which the device was intended to treat: _____

- 3) For each Coloplast Corp. pelvic mesh product identified above, indicate if, prior to implantation, you received any written and/or verbal information or instructions, including any risks or complications that might be associated with the use of the product(s)? **Yes** ____ **No** ____ **Don't Know** ____

If Yes:

- a. Provide the date you received the written and/or verbal information or instructions:

- b. Identify by name and address the person(s) who provided the information or instructions: _____

- c. What information or instructions did you receive? _____

d. If you have copies of the written information or instructions you received, please attach copies to your response.

4) For each Coloplast Corp. pelvic mesh product(s) that remains implanted in you:

a. Has any doctor recommended removal of the pelvic mesh product(s)?

Yes ____ **No** ____

If Yes, Identify by name and address the doctor who recommended removal and state your understanding of why the doctor recommended removal:

5) Have any of the Coloplast Corp., pelvic mesh product(s) been removed, in whole or in part?

Yes ____ **No** ____ **Don't Know** ____

If Yes, for each pelvic mesh product removed provide:

a. On what date, where and by whom (doctor) was the pelvic mesh product(s), or any portion of it, removed? _____

b. Explain why you consented to have the pelvic mesh product(s), or any portion of it, removed? _____

c. Does any medical treater, physician or anybody else on your behalf have possession of any portion of the pelvic mesh product® that was previously implanted in you and removed? **Yes** ____ **No** ____ **Don't Know** ____

If Yes, please state name and address of the person or entity having possession of same.

6) Do you claim that you suffered bodily injuries as a result of the implantation of any Coloplast Corp., pelvic mesh product(s)? **Yes** ____ **No** ____

If Yes:

a. Describe the bodily injuries, including any emotional or psychological injuries, that you claim resulted from the implantation of the pelvic mesh product(s).

- b. When is the first time you experienced symptoms of any of the bodily injuries you claim in your lawsuit to have resulted from the pelvic mesh product(s)?

- c. When did you first attribute these bodily injuries to the pelvic mesh product(s)?

- d. To the best of your knowledge and recollection, please state approximately when you first saw a health care provider for each of those bodily injuries you claim to have experienced relating to the pelvic mesh product(s):

- e. Are you currently experiencing symptoms related to your claimed bodily injuries?
Yes ____ **No** ____

If Yes, please describe your current symptoms in detail

- f. Are you currently seeing, or have you ever seen a doctor or healthcare provider for each of the bodily injuries or symptoms listed above? **Yes** ____ **No** ____

If Yes, please list all doctors you have seen for treatment of any of the bodily injuries you have listed above.

Provider Name and Address	Condition Treated	Approximate Dates of Treatment

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- g. Were you hospitalized at any time for the bodily injuries you listed above?
Yes ____ **No** ____

If Yes, please provide the following:

Hospital Name and Address	Condition Treated	Approximate Dates of Treatment

- 7) Other than the Coloplast Corp. pelvic mesh product(s) that are the subject of your lawsuit, have you been implanted with any other pelvic mesh products?
Yes ____ **No** ____

If Yes, please provide the following information:

- a. Product Name(s): _____

- b. Date of implantation procedure(s) and name and address of implanting doctor(s):

- c. Condition(s) sought to be treated through placement of the device(s):

- d. Whether the product(s) remain implanted inside of you today?
Yes ____ **No** ____

- 8) Are you making a claim for lost wages or lost earning capacity?
Yes ____ **No** ____

If Yes, state the annual gross income you derived from your employment for each year, beginning five years prior to the implantation of the pelvic mesh product(s) until the present:

- 9) Are you making a claim for lost out-of-pocket expenses?

Yes ____ **No** ____

If Yes, please identify and itemize all out-of-pocket expenses you have incurred:

- 10) Has anyone filed a loss of consortium claim in connection with your lawsuit regarding the pelvic mesh product(s)?

Yes ____ **No** ____

If Yes, identify by name and address the person who filed the loss of consortium claim, state the relationship of that person to you, and state the nature of the claim:

- 11) Please indicate whether the consortium plaintiff is alleging any of the claimed damages set forth below and itemize the alleged damages/expenses:

Claims	Yes/ No	Itemized Damages/Expenses
Loss of services of spouse		Not applicable
Impaired sexual relations		Not applicable
Lost wages/ lost earning capacity		
Lost out-of-pocket expenses		
Physical injuries		Not applicable
Psychological Injuries/ Emotional Injuries		Not applicable
Other		Not applicable

- 12) Please list the name and address of any healthcare providers the consortium plaintiff has seen for treatment for any physical, emotional, or psychological injuries or symptoms alleged to be related to the loss of consortium claim.

- 13) Have you or anyone acting on your behalf had any communication, oral or written, with any of the defendants or their representatives, other than your attorneys?

Yes ___ No ___ Don't Know ___

If Yes, set forth the date of the communication, the method of communication, the name of the person with whom you communicated, and the substance of the communication between you and any defendants or their representatives:

III. MEDICAL BACKGROUND

- 1) Provide your current age: _____ Height _____ Weight _____
- 2) At the time you received each pelvic mesh product(s), please state:
Your age _____ Your approximate weight _____
- 3) State number of vaginal births you have had? _____
- 4) State the number of cesarean section births you have had? _____
- 5) In chronological order, list any and all pelvic and/or abdominal surgeries you had at any time **BEFORE** implantation of the pelvic mesh product(s); identifying by name and address the doctor(s), hospital(s) or other healthcare provider(s) involved with each surgery; and providing the approximate date(s) for each. Insert additional rows as necessary.

Doctor or Healthcare Provider Involved (including address)	Description of Surgery Hospitalization	Approximate. Date

- 6) In chronological order, list any and all other surgeries, procedures, or hospitalizations you had in the 10 year period **BEFORE** implantation of the pelvic mesh product(s); identifying by name and address the doctor(s), hospital(s) or other healthcare provider(s) involved with each surgery or procedure; and providing the approximate date(s) for each. Insert additional rows as necessary.

Doctor or Healthcare Provider Involved (including address)	Description of Surgery Hospitalization	Approximate. Date

- 7) In chronological order, list any and all surgeries, procedures, or hospitalizations you had **AFTER** the implantation of the pelvic mesh product(s); identifying by name and address the doctor(s), hospital(s) or other healthcare provider(s) involved with each surgery or procedure; and provide the approximate date(s) for each. Insert additional rows as necessary.

Doctor or Healthcare Provider Involved (including address)	Description of Surgery/ Hospitalization	Approximate Date
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- 8) To the extent not already provided in the charts above, provide the name, address, and telephone number of every doctor, hospital, or other health care provider from which you have received medical advice and/or treatment for the past **10 years**. Insert additional rows as necessary.

Name and Specialty	Address	Approximate Dates/Years of Visits

- 9) Please describe your physical activities associated with daily living, physical fitness, household tasks, and employment-related activities *before* the implantation of each pelvic mesh product.

- 10) Please describe your physical activities associated with daily living, physical fitness, household tasks, and employment-related activities *after* the implantation of the pelvic mesh product(s).

- 11) To the best of your knowledge, have you suffered from any of the following:

Medical Condition		Sought treatment for?	Indicate whether condition occurred pre-implant, post-implant or both (explain, if necessary)
Adhesions	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Bleeding or Clotting Disorders If Yes, please specify disorder:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Bowel Obstruction	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Bowel Perforation	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Cancer If Yes, please specify type:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Chronic Constipation	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Collagen Disorder/Deficiency	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Connective Tissue Disorder If Yes, please specify disorder:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Crohn's Disease, Irritable Bowel Syndrome, Ulcerative Colitis, or Chronic Diarrhea	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____

If Yes , please specify which condition and treatment prescribed:			
Cystocele	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Diabetes	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Diverticulitis	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Dyspareunia	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Enterocoele	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Fistulas	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Hernias	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Hypertension or High Blood Pressure	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Hypotension or Low Blood Pressure	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Immune System Disease or Dysfunction including HIV/AIDS If Yes , please specify condition:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Malnutrition	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Muscle or Muscle-Wasting Disorder If Yes , please specify disorder:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Neuromuscular Disease or Disorder	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____

If Yes , please specify disorder:			
Obesity	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Pelvic Trauma If Yes , please describe trauma:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Pelvic Tumors or Fibroids	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Peritonitis/Sepsis	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Rectocele	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Recurrent or Chronic Vaginal or Bladder Infections If Yes , please specify location and nature of infections:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Recurrent Vaginal Pain If Yes , please describe the nature of pain experienced:	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Urinary Incontinence	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Urinary Retention	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Uterine Prolapse	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Vaginal Vault Prolapse	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____
Wound Healing Problems	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____

If Yes , please explain:			
Any other disease of the gut, intestines, or bowels If Yes , please specify condition (s):	Yes ____ No ____	Yes ____ No ____	Pre ____ Post ____

THE FOLLOWING QUESTIONS ARE CONFIDENTIAL AND SUBJECT TO THE PROTECTIVE ORDER APPLICABLE TO THIS CASE.

- a) **Were you diagnosed with and/or treated for Sexually Transmitted Diseases for the five year period prior to the implantation of the pelvic mesh product(s) through the present?**
- Yes ____ No ____

If Yes, specify the disease, date of onset, medication/treatment, treating physician and current status of condition:

- b) **Have you been diagnosed with and/or treated for any alcohol or chemical dependency for the one year prior to the implantation of the pelvic mesh product(s) through the present?**
- Yes ____ No ____

If Yes, specify type and time period of dependency, type of treatment received, name of treatment provider, and current status of condition:

- c) **Have you experienced, been diagnosed with or been treated for any mental health conditions including depression, anxiety or other emotional or psychiatric disorders in the 5 year period before implantation of the pelvic mesh product(s) through the present?**
- Yes ____ No ____

If Yes, specify condition, date of onset, medication/treatment, treating physician and current status of condition:

12) Have you experienced menopause? Yes ____ No ____

If Yes, at what age did it begin? _____

13) Have you undergone vaginal estrogen therapy, hormone therapy, or systemic estrogen replacement therapy (ERT)? Yes ____ No ____

If Yes, please provide the type of therapy you received, date(s) of the therapy, and the name and address of the healthcare provider providing the therapy.

14) Do you now or have you ever smoked tobacco products? Yes ____ No ____

If Yes:

a) How long have/did you smoke, and how many packs per day did you smoke?

15) List each prescription medication you have taken **for more than 3 months at a time, within the last 5 years prior to implant to present**, giving the name and address of the pharmacy where you received/filled the medication, the reason you took the medication, and the approximate dates of use.

Medication and Dosage	Pharmacy (Name and Address)

IV. INSURANCE INFORMATION

- 1) Provide the following information for any past or present medical insurance coverage within the last 10 years:

Insurance Company (Name and Address)	Policy Number	Name of Policy Holder/Insured (if different than you)	Approx. Dates of Coverage

- 2) Have you ever been denied life insurance for reasons relating to your health?

Yes ____ **No** ____ **Don't Know** ____

If Yes, please state when the denial occurred, the name of the life insurance company, and the company's reason for denial: _____

- 3) To the best of your knowledge, have you been approved to receive or are you receiving Medicare benefits due to age, disability, condition or any other reason or basis?

Yes ____ **No** ____

If Yes, please specify the following:

- a) The date on which you first became eligible: _____

[Please note: if you are not currently a Medicare-eligible beneficiary, but become eligible for Medicare during the pendency of this lawsuit, you must supplement your response at that time. This information is necessary for all parties to comply with Medicare regulations. See 42 U.S.C. 1395y(b)(8), also known as Section 111 of the Medicare, Medicaid and SCHIP Extension Act of 2007 and 42 U.S.C. 1395y(b)(2) also known as the Medicare Secondary Payer Act.]

V. PRIOR CLAIM INFORMATION

- 1) Have you filed a lawsuit or made a claim in the last 10 years, other than in the present suit relating to any bodily injury?

Yes ____ **No** ____

If Yes, please specify the following:

- a) Court in which suit/claim filed or made: _____
- b) Case/Claim Number: _____
- c) Nature of Claim/Injury: _____

- 2) Have you applied for workers' compensation (WC), Social Security disability (SSI or SSD) benefits, or other state or federal disability benefits within the past 10 years?

Yes ____ **No** ____

If Yes, please specify the following:

- a) Date (or year) of application: _____
 - b) Type of benefits sought _____
 - c) Agency/Insurer from which you sought the benefits: _____
 - d) The nature of the claimed injury/disability: _____
 - e) Whether the claim was accepted or denied: _____
- _____

VI. FACT WITNESSES

- 1) Please identify all persons who you believe possess information concerning your injury(ies) and current medical conditions, other than your healthcare providers, and please state their name address and his/her/their relationship to you:

Name	Address	Relationship to You	Information you Believe Person Possesses

VII. IDENTIFICATION OF DOCUMENTS AND OTHER ELECTRONICALLY STORED INFORMATION

For the period beginning three years prior to implantation of the pelvic mesh product(s) to present, please identify all research, including on-line research, you have conducted regarding the subjects of this litigation, including the implantation of the pelvic mesh product(s), the injuries and/or damages you claim resulted from the implantation of the pelvic mesh product(s), or your medical or physical condition. Identify date, time, and source, including any websites visited. Research conducted to understand the legal and strategic advice of your counsel is not considered responsive to this request.

VIII. DOCUMENT REQUESTS

1) **RELEASES.**

NOTE: Please sign and attach to this Fact Sheet the authorizations for the release of records appended hereto.

2) **DOCUMENTS.** State whether you have any of the following documents in your possession, custody, and/or control. If you do, please provide a true and correct copy of any such documents with this completed Fact Sheet.

- a) If you were appointed by a court to represent the plaintiff in this lawsuit, produce any documents demonstrating your appointment as such.
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
- b) If you represent the estate of a deceased person in this lawsuit, produce a copy of the decedent's death certificate and autopsy report (if applicable).
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
- c) Produce any communications (sent or received) in your possession, which shall include materials accessible to you from any computer on which you have sent or received such communications, concerning the pelvic mesh product(s) or subject litigation, including but not limited to all letters, e-mails, blogs, Facebook posts, tweets, newsletters, etc. sent or received by you. Research conducted to understand

the legal and strategic advice of your counsel is not considered responsive to this request.

i. Not Applicable _____

ii. The documents are attached _____ [OR] I have no documents _____

- d) Produce all documents (including journal entries, lists, memoranda, notes, diaries), photographs, video, DVDS or other media, including all copies, discussing or referencing the subjects of this litigation including the pelvic mesh product(s), the injuries and/or damages you claim resulted from the pelvic mesh product(s), or evidencing your physical condition from three years prior to the implantation of the pelvic mesh product(s) to present, including but not limited to the injuries for which you claim relief in this lawsuit. Research conducted to understand the legal and strategic advice of your counsel is not considered responsive to this request.

i. Not Applicable _____

ii. The documents are attached _____ [OR] I have no documents _____

- e) Produce any pelvic mesh product packaging, labeling, advertising, or any other pelvic mesh product product-related items in your possession, custody or control.

i. Not Applicable _____

ii. The documents are attached _____ [OR] I have no documents _____

- f) Produce all documents concerning any communication between you and the Food and Drug Administration (FDA) or between you and any employee or agent of the Defendants, regarding the pelvic mesh product(s) at issue, except as to those communications which are attorney client/work product privileged.

i. Not Applicable _____

ii. The documents are attached _____ [OR] I have no documents _____

- g) Produce all documents in your possession, custody or control evidencing or relating to any correspondence or communication between either you or Coloplast Corp., (or any of its related companies or divisions) with any of your doctors, healthcare providers, and/or you relating to the pelvic mesh product(s), except as to those communications which are attorney client/work product privileged.

- i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
 - h) Produce any and all documents in your possession, custody or control reflecting, describing, or in any way relating to any instructions or warnings you received prior to implantation of any pelvic mesh product(s) concerning the risks and/or benefits of your surgery, including but not limited to any risks and/or benefits associated with the pelvic mesh product(s).
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
 - i) Produce any and all documents reflecting the model number and lot number of the pelvic mesh product(s) you received.
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
 - j) If you underwent surgery to explant in whole or in part the pelvic mesh product(s) that you received: produce any and all documents in your possession, custody or control aside from documents that may have been generated by experts retained by your counsel for litigation purposes, relating to any evaluation of the pelvic mesh product(s) and any other material that was (were) surgically removed from you.
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
 - k) If you claim lost wages or lost earning capacity, copies of your federal and state tax returns for the two years prior to implantation of the pelvic mesh product(s) to the present.
 - i. Not Applicable _____
 - ii. The documents are attached _____ [OR] I have no documents _____
- If you do not have the required tax returns, you must provide fully completed authorizations for the release of federal and state tax returns.
- l) All documents in your possession, custody or control concerning payment by Medicare on the injured party's behalf relating to the injuries claimed in this lawsuit, including but not limited to Interim Conditional Payment summaries and/or estimates prepared by Medicare or its representatives regarding payments made on

your behalf for medical expenses relating to the subject of this litigation.

i. Not Applicable _____

ii. The documents are attached _____ [OR] I have no documents _____

[Please note: if you are not currently a Medicare-eligible beneficiary, but become eligible for Medicare during the pendency of this lawsuit, you must supplement your response at that time. This information is necessary for all parties to comply with Medicare regulations. See 42 U.S.C. 1395y(b)(8), also known as Section 111 of the Medicare, Medicaid and SCHIP Extension Act of 2007 and 42 U.S.C. 1395y(b)(2) also known as the Medicare Secondary Payer Act.]

VERIFICATION

I, _____, declare under penalty of perjury subject to all applicable laws, that I have carefully reviewed the final copy of this Plaintiff Fact Sheet dated _____ and verified that all of the information provided is true and correct to the best of my knowledge, information and belief.

Signature of Plaintiff

VERIFICATION OF LOSS OF CONSORTIUM

I, _____, declare under penalty of perjury subject to all applicable laws, that I have carefully reviewed the final copy of this Plaintiff Fact Sheet dated _____ and verified that all of the information provided is true and correct to the best of my knowledge, information and belief.

Signature of Consortium Plaintiff

APPENDIX "A"
(Authorization Forms)

Instructions for Using this Form

Complete this form only if you want us to give information or records about you, a minor, or a legally incompetent adult, to an individual or group (for example, a doctor or an insurance company). If you are the natural or adoptive parent or legal guardian, acting on behalf of a minor child, you may complete this form to release only the minor's non-medical records. We may charge a fee for providing information unrelated to the administration of a program under the Social Security Act.

NOTE: Do not use this form to:

- Request the release of medical records on behalf of a minor child. Instead, visit your local Social Security office or call our toll-free number, 1-800-772-1213 (TTY-1-800-325-0778), or
- Request detailed information about your earnings or employment history. Instead, complete and mail form SSA-7050-F4. You can obtain form SSA-7050-F4 from your local Social Security office or online at www.ssa.gov/online/ssa-7050.pdf.

How to Complete this Form

We will not honor this form unless all required fields are completed. An asterisk (*) indicates a required field. Also, we will not honor blanket requests for "any and all records" or the "entire file." You must specify the information you are requesting and you must sign and date this form. We may charge a fee to release information for non-program purposes.

- Fill in your name, date of birth, and social security number or the name, date of birth, and social security number of the person to whom the requested information pertains.
- Fill in the name and address of the person or organization where you want us to send the requested information.
- Specify the reason you want us to release the information.
- Check the box next to the type(s) of information you want us to release including the date ranges, where applicable.
- You, the parent or the legal guardian acting on behalf of a minor child or legally incompetent adult, must sign and date this form and provide a daytime phone number.
- If you are not the individual to whom the requested information pertains, state your relationship to that person. We may require proof of relationship.

PRIVACY ACT STATEMENT

Section 205(a) of the Social Security Act, as amended, authorizes us to collect the information requested on this form. We will use the information you provide to respond to your request for access to the records we maintain about you or to process your request to release your records to a third party. You do not have to provide the requested information. Your response is voluntary; however, we cannot honor your request to release information or records about you to another person or organization without your consent. We rarely use the information provided on this form for any purpose other than to respond to requests for SSA records information. However, the Privacy Act (5 U.S.C. § 552a(b)) permits us to disclose the information you provide on this form in accordance with approved routine uses, which include but are not limited to the following:

- 1.To enable an agency or third party to assist Social Security in establishing rights to Social Security benefits and or coverage;
- 2.To make determinations for eligibility in similar health and income maintenance programs at the Federal, State, and local level;
- 3.To comply with Federal laws requiring the disclosure of the information from our records; and,
- 4.To facilitate statistical research, audit, or investigative activities necessary to assure the integrity of SSA programs.

We may also use the information you provide when we match records by computer. Computer matching programs compare our records with those of other Federal, State, or local government agencies. We use information from these matching programs to establish or verify a person's eligibility for Federally-funded or administered benefit programs and for repayment of incorrect payments or overpayments under these programs. Additional information regarding this form, routine uses of information, and other Social Security programs is available on our Internet website, www.socialsecurity.gov, or at your local Social Security office.

PAPERWORK REDUCTION ACT STATEMENT

This information collection meets the requirements of 44 U.S.C. § 3507, as amended by section 2 of the Paperwork Reduction Act of 1995. You do not need to answer these questions unless we display a valid Office of Management and Budget control number. We estimate that it will take about 3 minutes to read the instructions, gather the facts, and answer the questions. **SEND OR BRING THE COMPLETED FORM TO YOUR LOCAL SOCIAL SECURITY OFFICE. You can find your local Social Security office through SSA's website at www.socialsecurity.gov. Offices are also listed under U.S. Government agencies in your telephone directory or you may call 1-800-772-1213 (TTY 1-800-325-0778).** You may send comments on our time estimate above to: SSA, 6401 Security Blvd., Baltimore, MD 21235-6401. ***Send only comments relating to our time estimate to this address, not the completed form.***

Consent for Release of Information

You must complete all required fields. We will not honor your request unless all required fields are completed. (**signifies a required field*).

TO: Social Security Administration

***My Full Name**

***My Date of Birth**
(MM/DD/YYYY)

***My Social Security Number**

I authorize the Social Security Administration to release information or records about me to:

***NAME OF PERSON OR ORGANIZATION:**

***ADDRESS OF PERSON OR ORGANIZATION:**

***I want this information released because:**

We may charge a fee to release information for non-program purposes.

***Please release the following information selected from the list below:**

You must specify the records you are requesting by checking at least one box. We will not honor a request for "any and all records" or "my entire file." Also, we will not disclose records unless you include the applicable date ranges where requested.

1. ☐ Social Security Number
2. ☐ Current monthly Social Security benefit amount
3. ☐ Current monthly Supplemental Security Income payment amount
4. ☐ My benefit or payment amounts from date _____ to date _____
5. ☐ My Medicare entitlement from date _____ to date _____
6. ☐ Medical records from my claims folder(s) from date _____ to date _____

If you want us to release a minor child's medical records, do not use this form. Instead, contact your local Social Security office.

7. ☐ Complete medical records from my claims folder(s)
8. ☐ Other record(s) from my file (**you must specify the records you are requesting, e.g., doctor report, application, determination or questionnaire**)

I am the individual, to whom the requested information or record applies, or the parent or legal guardian of a minor, or the legal guardian of a legally incompetent adult. I declare under penalty of perjury (28 CFR § 16.41(d)(2004)) that I have examined all the information on this form, and any accompanying statements or forms, and it is true and correct to the best of my knowledge. I understand that anyone who knowingly or willfully seeks or obtain access to records about another person under false pretenses is punishable by a fine of up to \$5,000. I also understand that I must pay all applicable fees for requesting information for a non-program-related purpose.

***Signature:** _____ ***Date:** _____

***Address:** _____

Relationship (if not the subject of the record): _____ ***Daytime Phone:** _____

Witnesses must sign this form ONLY if the above signature is by mark (X). If signed by mark (X), two witnesses to the signing who know the signee must sign below and provide their full addresses. Please print the signee's name next to the mark (X) on the signature line above.

1. Signature of witness

2. Signature of witness

Address(Number and street, City, State, and Zip Code)

Address(Number and street, City, State, and Zip Code)



Medicare

Beneficiary Services: 1-800-MEDICARE (1-800-633-4227)
TTY/TDD: 1-877-486-2048

This form is used to advise Medicare of the person or persons you have chosen to have access to your personal health information.

Where to Return Your Completed Authorization Forms:

After you complete and sign the authorization form, return it to the address below:

Medicare BCC, Written Authorization Dept.
PO Box 1270
Lawrence, KS 66044

For New York Medicare Beneficiaries ONLY

The New York State Public Health Law protects information that reasonably could identify someone as having HIV symptoms or infection, and information regarding a person's contacts. Because of New York's laws protecting the privacy of information related to alcohol and drug abuse, mental health treatment, and HIV, there are special instructions for how you, as a New York resident, should complete this form.

- For question 2A, check the box for *Limited Information*, even if you want to authorize Medicare to release any and all of your personal health information.
- **Then proceed to question 2B.** You may also check any of the remaining boxes and include any additional limitations in the space provided. For example, you could write "payment information".

Medicare BCC, Written Authorization Dept.
PO Box 1270
Lawrence, KS 66044

Instructions for Completing Section 2C of the Authorization Form:

Please select one of the following options.

- **Option 1** To **include** all information, check the box: "all information, including information about alcohol and drug abuse, mental health treatment, and HIV". Proceed with the rest of the form.
- **Option 2** To **exclude** the information listed above, check the box: "Exclude information about alcohol and drug abuse, mental health treatment and HIV". Then proceed with the rest of the form.

If you have any questions or need additional assistance, please feel free to call us at 1-800-MEDICARE (1-800-633-4227). TTY users should call 1-877-486-2048.

Sincerely,

1-800-MEDICARE
Customer Service Representative

Encl.

Information to Help You Fill Out the “1-800-MEDICARE Authorization to Disclose Personal Health Information” Form

By law, Medicare must have your written permission (an “authorization”) to use or give out your personal medical information for any purpose that isn't set out in the privacy notice contained in the Medicare & You handbook. You may take back (“revoke”) your written permission at any time, except if Medicare has already acted based on your permission.

If you want 1-800-MEDICARE to give your personal health information to someone other than you, you need to let Medicare know in writing.

If you are requesting personal health information for a deceased beneficiary, please include a copy of the legal documentation which indicates your authority to make a request for information. (For example: Executor/Executrix papers, next of kin attested by court documents with a court stamp and a judge's signature, a Letter of Testamentary or Administration with a court stamp and judge's signature, or personal representative papers with a court stamp and judge's signature.) Also, please explain your relationship to the beneficiary.

Please use this step by step instruction sheet when completing your “1-800-MEDICARE Authorization to Disclose Personal Health Information” Form. Be sure to complete all sections of the form to ensure timely processing.

1. Print the name of the person with Medicare.

Print the Medicare number exactly as it is shown on the red, white, and blue Medicare card, including any letters (for example, 000000000A).

Print the birthday in month, day, and year (mm/dd/yyyy) of the person with Medicare.

- 2.** This section tells Medicare what personal health information to give out. Please check a box in 2A to indicate how much information Medicare can disclose. If you only want Medicare to give out limited information (for example, Medicare eligibility), also check the box(es) in 2B that apply to the type of information you want Medicare to give out. Box 2C must be completed by **New York Residents**.
- 3.** This section tells Medicare when to start and/or when to stop giving out your personal health information. Check the box that applies and fill in dates, if necessary.
- 4.** Medicare will give your personal health information to the person(s) or organization(s) you fill in here. You may fill in more than one person or organization.

If you designate an organization, you must also identify one or more individuals in that organization to whom Medicare may disclose your personal health information.

5. The person with Medicare or personal representative must sign their name, fill in the date, and provide the phone number and address of the person with Medicare.

If you are a personal representative of the person with Medicare, check the box, provide your address and phone number, and attach a copy of the paperwork that shows you can act for that person (for example, Power of Attorney).

6. Send your completed, signed authorization to Medicare at the address shown here on your authorization form.
7. If you change your mind and don't want Medicare to give out your personal health information, write to the address shown under number six on the authorization form and tell Medicare. Your letter will revoke your authorization and Medicare will no longer give out your personal health information (except for the personal health information Medicare has already given out based on your permission).

You should make a copy of your signed authorization for your records before mailing it to Medicare.

1-800-MEDICARE Authorization to Disclose Personal Health Information

Use this form if you want 1-800-MEDICARE to give your personal health information to someone other than you.

- | | | |
|---|---|--------------------------------------|
| 1. Print Name
(First and last name of the person with Medicare) | Medicare Number
(Exactly as shown on the Medicare Card) | Date of Birth
(mm/dd/yyyy) |
|---|---|--------------------------------------|

2. Medicare will only disclose the personal health information you want disclosed.

2A: Check only one box below to tell Medicare the specific personal health information you want disclosed:

- ☐ Limited Information (go to question 2b)
- ☐ Any Information (go to question 3)

2B: Complete only if you selected “limited information”. Check all that apply:

- ☐ Information about your Medicare eligibility
- ☐ Information about your Medicare claims
- ☐ Information about plan enrollment (e.g. drug or MA Plan)
- ☐ Information about premium payments
- ☐ Other Specific Information (please write below; for example, payment information)

2C: NY Residents Only, this section must be completed.

Please select one of the following options: (Please check only one box.)

- ☐ Include all information. This includes information about alcohol and drug abuse, mental health treatment, and HIV.

OR

- ☐ Exclude information about alcohol and drug abuse, mental health treatment, and HIV.

- 3. Check only one box below indicating how long Medicare can use this authorization to disclose your personal health information** (subject to applicable law—for example, your State may limit how long Medicare may give out your personal health information):

☐ Disclose my personal health information indefinitely

☐ Disclose my personal health information for a specified period only

beginning: _____ (mm/dd/yyyy) and ending: _____ (mm/dd/yyyy)

- 4. Fill in the name and address of the person or organization to whom you want Medicare to disclose your personal health information. Please provide the specific name of the person for any organization you list below. If you would like to authorize any additional individuals or organizations, please add those to the back of this form.**

Name Norton Rose Fulbright US LLP

Address 98 San Jacinto Boulevard, STE 1100, Austin, TX 78701

Name Eastwood Nurse Legal Consulting LLC

Address 200 Lake Rd., Bozeman, MT 59718-8522

Note: You have the right to take back (“revoke”) your authorization at any time, in writing, except to the extent that Medicare has already acted based on your permission. To revoke authorization, send a written request to the address noted below. Your authorization or refusal to authorize disclosure of your personal health information will have no effect on your enrollment, eligibility for benefits, or the amount Medicare pays for the health services you receive.

5.

I authorize 1-800-MEDICARE to disclose my personal health information listed above to the person(s) or organization(s) I have named on this form. I understand that my personal health information may be re-disclosed by the person(s) or organization(s) and may no longer be protected by law.

Signature

Telephone Number

Date (mm/dd/yyyy)

Print the address of the person with Medicare (Street Address, City, State, and ZIP)

☐ Check here if you are signing as a personal representative and complete below.
Please attach the appropriate documentation (for example, Power of Attorney). This only applies if someone other than the person with Medicare signed above.

Print the Personal Representative's Address (Street Address, City, State, and ZIP)

Telephone Number of Personal Representative: _____

Personal Representative's Relationship to the Beneficiary: _____

6. Send the completed, signed authorization to:

Medicare BCC, Written Authorization Dept.
PO Box 1270
Lawrence, KS 66044

PrintForm

Note: You have the right to take back (“revoke”) your authorization at any time, in writing, except to the extent that Medicare has already acted based on your permission. If you would like to revoke authorization, send a written request to the address noted above.

Your authorization or refusal to authorize disclosure of your personal health information will have no effect on your enrollment, eligibility for benefits, or the amount Medicare pays for the health services you receive.

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0938-0930. The time required to complete this information collection is estimated to average 15 minutes per response, including the time to review instructions, search existing data resources, gather the data needed, and complete and review the information collection. If you have comments concerning the accuracy of the time estimate(s) or suggestions for improving this form, please write to: CMS, 7500 Security Boulevard, Attn: PRA Reports Clearance Officer, Mail Stop C4-26-05, Baltimore, Maryland 21244-1850.

**IN THE UNITED STATES DISTRICT COURT
FOR THE SOUTHERN DISTRICT OF WEST VIRGINIA
CHARLESTON DIVISION**

IN RE: COLOPLAST CORP., PELVIC
SUPPORT SYSTEMS PRODUCTS
LIABILITY LITIGATION

MDL 2387

THIS DOCUMENT RELATES TO

Civil Action No.: _____

Name of Plaintiff

DEFENDANT FACT SHEET

For each case involving a Coloplast product, Defendant Coloplast Corp. (hereinafter "Coloplast" or "Defendant") shall produce any non-privileged matter that is relevant to the request pursuant to Rule 26(b) of the Federal Rules of Civil Procedure. Except as otherwise set forth in any Order, this Fact Sheet must be completed and served on Plaintiffs' counsel in each individual case by _____.

I. INFORMATION

Case Caption: _____

Case Number: _____

II. IMPLANTING AND EVALUATING PHYSICIANS

Plaintiff has identified physicians in Sections II.1 ("Implanting Physician") and II.6.f. of the Plaintiff's Fact Sheet (together, "Identified Physicians"). As to each Identified Physician, provide or produce the following information:

A. NON-SALES REPRESENTATIVE CONSULTATION CONTACTS

As to each Identified Physician with whom the Defendant was affiliated or consulted regarding the female pelvic mesh product(s) (outside the context of sales representative contacts), Defendant will provide the following for each Identified Physician:

1. Identify the Identified Physician.
2. Any communications between the Defendant and the Identified Physician relevant to Defendant's female pelvic mesh product(s), including communications regarding the safety, use, or efficacy of Defendant's female pelvic mesh products.

3. Any monetary or non-monetary benefits provided to the Identified Physician by the Defendant.
4. Written agreements, including contracts, setting forth the terms or nature of any consultation or affiliation with the Identified Physician related to Defendant's female pelvic mesh product(s); this includes but is not limited to any contracts to research or otherwise study Defendant's female pelvic mesh product(s).
5. For each facility identified by the Plaintiff as one in which she had a surgical implant of Defendant's product by the Implanting Physician, set forth the number and type of women's pelvic mesh product(s) purchased from Defendant for five (5) years prior to the date of surgery.
6. Set forth any communication between the Defendant and the Identified Physician with regard to the Plaintiff.

B. SALES REPRESENTATIVES

For each Implanting Physician identified by Plaintiff in the Plaintiff Fact Sheet, set forth the following or produce relevant, non-privileged documents:

1. Identify the Implanting Physician.
2. Identity of Defendant's sales representative(s), if any, that have had contact with the Implanting Physician regarding Defendant's female pelvic mesh product(s).
3. State whether or not the Defendant's sales representative(s) identified above is currently employed by Defendant. If the sales representative(s) is no longer employed by Defendant, please provide the last known address of the sales representative(s).
4. The name and model number for the female pelvic mesh product(s) that the sales representative was responsible for with regard to the Implanting Physician.
5. Identify all district and/or regional sales managers who were responsible for the management of the sales representatives identified in your response to Number 2 above, and their current relationship, if any, with Defendant.
6. Set forth all information related to the use of Defendant's female pelvic mesh product(s) provided by the physician to the sales representative, with regard to the Plaintiff.
7. Identify any reprimand or rebuke for any non-privileged relevant reason of the sales representative while employed by you.

III. INFORMATION REGARDING THE PLAINTIFF

- A. Outside of information exchanged as part of this litigation, identify all data, information, objects, and reports in Defendant's possession or control specific to the Plaintiff. Attorney-work product is specifically excluded.
- B. Identify any contact, either written or oral, between the Plaintiff and any employee or representative of the Defendant, including but not limited to pre-operative inquiries, and post-operative complaints. This request specifically includes, but is not limited to, calls to your Customer Care Center.
- C. Identify all Med Watch Adverse Event Reports and/or any other documents submitted to the FDA or any other government agency with regard to the Plaintiff.
- D. If Plaintiff has provided a lot number for Defendant's product in Section II.1 of the Plaintiff's Fact Sheet, state whether such lot number has ever been subject to a recall.
- E. Identify all communications that Defendant has had with any of the Plaintiff's Identified Physicians specifically regarding the Plaintiff.
- F. If you have any evidence or records indicating or demonstrating the possibility that any person, entity, condition, or product, other than the Defendant and its product(s), is a cause of the Plaintiff's injuries, ("Alternate Cause") set forth:
 - 1. Identify the Alternate Cause with specificity.
 - 2. Set forth the date and mechanism of alternate causation.

IV. MANUFACTURING INFORMATION

- A. With respect to any Defendant's female pelvic mesh product implanted in the Plaintiff and which is the subject of this suit (where Plaintiff has provided a lot number for Defendant's product in Section II.1 of the Plaintiff's Fact Sheet), identify the device history record.
- B. Where Plaintiff has provided a lot number for Defendant's product in Section II.1 of the Plaintiff's Fact Sheet and has identified the implanting facility, identify the date of order, invoicing and shipping, and the person or entity purchasing, for each of Plaintiff's implanted device(s).
- C. Identify manufacturers of Defendant's female pelvic mesh devices

implanted in the Plaintiff.

V. DOCUMENTS

Please provide the following documents pursuant to an ESI protocol to be agreed upon by the parties or entered by the Court.

- A. The specific documentation described in I through IV above; except you need not serve copies of medical records that were provided to Defendant by Plaintiff's counsel.
- B. The specific documentation described in section II above; except you need not serve copies of medical records that were provided to Defendant by Plaintiff's counsel.
- C. The specific documentation described in section III above; except you need not serve copies of medical records that were provided to Defendant by Plaintiff's counsel.
- D. The specific documentation described in section IV above; except you need not serve copies of medical records that were provided to Defendant by Plaintiff's counsel.
- E. Aside from any privileged or attorney-work product materials, produce all documents that refer, reference, attribute, or allude to the Plaintiff in Defendant's possession or control, to the extent not identified and attached in response to a prior question.
- F. For a lot number identified by Plaintiff in Section II.1 of Plaintiff's Fact Sheet, produce a true and complete copy of the device history record.
- G. Produce a true and complete copy of the complaint file relating to the Plaintiff.
- H. All call notes, details notes, or call summaries made by Defendant's sales representatives regarding each Identified Physician relating to Defendant's female pelvic mesh product(s) during the relevant time period.
- I. Any communications between Defendant or its sales representatives and the Identified Physician regarding Defendant's female pelvic mesh product(s).

VERIFICATION

I am an authorized agent of Coloplast and I verify the Defendant's Response to Defendant's Fact Sheet in *In Re Coloplast Corp., Pelvic Support Systems Products Liability Litigation*, MDL No. 2387 (S.D. W. Va.). The matters stated therein are not my personal knowledge; the facts stated herein have been assembled by authorized employees and counsel of Coloplast Corp. and I am informed that the facts stated herein are true. I hereby certify, in my authorized capacity as an agent for Coloplast Corp., that the responses to the aforementioned Defendant's Fact Sheet are true and complete to the best of Coloplast Corp.'s knowledge.

Date: _____

Coloplast Corp.