IN THE UNITED STATES DISTRICT COURT FOR THE MIDDLE DISTRICT OF TENNESSEE NASHVILLE DIVISION

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COMPLAINT

Plaintiff Amanda Scott, Individually and as Representative of the Estate of Ray Herndon Celsor, deceased, (herein after "Plaintiff"), by and through Plaintiff's attorneys, brings this action for personal injuries and wrongful death against Defendants Boehringer Ingelheim Pharmaceuticals, Inc., Boehringer Ingelheim Corporation, Boehringer Ingelheim USA Corporation, Boehringer Ingelheim Vetmedica, Boehringer Ingelheim Pharma GMBH & Co. KG, and Boehringer Ingelheim International GMBH (collectively, "Boehringer Ingelheim" or "Defendants"). Plaintiff alleges as follows:

PARTIES

- 1. Plaintiff Amanda Scott is the adult child of Ray Herndon Celsor, deceased, and the Representative of the Estate of Ray Herndon Celsor. She is bringing her individual claims, including her claim for the wrongful death of Ray Herndon Celsor, and the claims of the estate. At all times relevant hereto, Plaintiff, Amanda Scott, was a resident and citizen of Clarksville, Montgomery County, Tennessee. Ray Herndon Celsor will be referred to herein as Ray Herndon Celsor and/or Plaintiff.
- 2. Boehringer Ingelheim Pharmaceuticals, Inc. ("Boehringer") is a Delaware corporation which has its principal place of business at 900 Ridgebury Road, Ridgefield, Connecticut 06877. Boehringer has conducted business and derived substantial revenue from within the State of Tennessee, and may be served through its registered agent for service of process, CT Corporation System, 800 S. Gay Street, Ste. 2021, Knoxville, Tennessee 37929-9710.
- 3. Boehringer Ingelheim Vetmedica, Inc. ("Boehringer Vet") is a Delaware corporation which has its principal place of business at 2621 North Belt Highway, St. Joseph, MO 64506. Boehringer Vet has conducted business and derived substantial revenue from within the State of Tennessee, and may be served through its registered agent for service of process, CT Corporation System, 800 S. Gay Street, Ste. 2021, Knoxville, Tennessee 37929-9710.
- 4. Boehringer Ingelheim Corporation ("Boehringer Co") is a Delaware corporation which has its principal place of business at 900 Ridgebury Road, Ridgefield, Connecticut 06877. Boehringer Co. has conducted business and derived substantial

revenue from within the State of Tennessee, but does not maintain a regular place of business in this state or a designated agent for service of process for proceedings that arise out of Boehringer Co's business done in this state. Boehringer Co may be served with citation by its registered agent for service of process, CT Corporation System, One Corporate Center, Hartford, CT 06103.

- 5. Boehringer Ingelheim USA Corporation ("Boehringer USA") is a Delaware corporation which has its principal place of business at 900 Ridgebury Road, Ridgefield, Connecticut 06877. Boehringer USA has conducted business and derived substantial revenue from within the State of Tennessee, but does not maintain a regular place of business in this state or a designated agent for service of process for proceedings that arise out of Boehringer USA's business done in this state. Boehringer USA may be served with citation by its registered agent for service of process, CT Corporation System, One Corporate Center, Hartford, CT 06103.
- 6. Boehringer Ingelheim Pharma GmbH & Co. KG ("Boehringer Pharma") is a foreign corporation with its principal place of business located at Boehringer Ingelheim Pharma GmbH & Co. KG, Binger Strasse 173, 55216 Ingelheim am Rhein, Germany. Boehringer Pharma has transacted and conducted business within the State of Tennessee. Boehringer Pharma has derived substantial revenue from goods and products disseminated and used in the State of Tennessee, and Boehringer Pharma expected or should have expected their acts to have consequences within the State of Tennessee, and derived substantial revenue from commerce within the State of Tennessee.

7. Boehringer Ingelheim International GmbH ("Boehringer International") is a foreign corporation with its principal place of business located at Boehringer Ingelheim International GmbH, Binger Strasse 173, 55216 Ingelheim am Rhein, Germany. Boehringer International has transacted and conducted business within the State of Tennessee. Boehringer International has derived substantial revenue from goods and products disseminated and used in the State of Tennessee, and Boehringer International expected or should have expected their acts to have consequences within the State of Tennessee, and derived substantial revenue from commerce within the State of Tennessee.

JURISDICTION AND VENUE

- 8. Jurisdiction is proper in this court pursuant to 28 USC §1332 for the reason that there is complete diversity of citizenship between Plaintiffs and Defendants and the matter in controversy greatly exceeds the sum of seventy-five thousand dollars (\$75,000.00), exclusive of interest and costs.
- 9. This Court has jurisdiction over the non-resident Defendants because they have done business in the State of Tennessee, have committed a tort in whole or in part in the State of Tennessee, and have continuing contacts with the State of Tennessee.
- 10. Venue of this case is proper in the Middle District of Tennessee pursuant to 28 U.S.C. § 1391(b)(1) because Defendants are residents of this state.

11. Venue is further proper in this Court pursuant to 28 U.S.C. § 1391 because a substantial part of the events giving rise to Plaintiff's claims occurred, in part, in the Middle District of Tennessee.

FACTUAL BACKGROUND

Background of the Case

- 12. At all relevant times, Defendants, directly or through their agents, apparent agents, servants or employees designed, manufactured, marketed, advertised, distributed, promoted, labeled, tested and sold Pradaxa® (dabigatran etexilate mesylate).
- 13. Pradaxa® is a direct thrombin inhibitor that is indicated to reduce the risk of stroke and systemic embolism in patients with non-valvular atrial fibrillation. Patients with atrial fibrillation have an increased risk of stroke.
- 14. Pradaxa® was approved by the Food and Drug Administration ("FDA") on October 19, 2010. The FDA approved two dosages: 75 mg and 150 mg, to be taken twice daily. Pradaxa® was the first anticoagulation medication approved in the U.S. in more than 50 years for patients with non-valvular atrial fibrillation.
- 15. Prior to the FDA's approval of Pradaxa®, warfarin was the only oral anticoagulation available in the U.S. for reducing stroke and systemic embolism in patients with atrial fibrillation. Unlike patients who use Pradaxa®, users of warfarin must follow dietary restrictions and regularly monitor their blood levels (INR) by undergoing blood tests and potentially adjusting the dose of their medication.

<u>Defendants' over promotion of Pradaxa®</u>

- 16. Defendants promoted Pradaxa® as a novel medicine for patients with non-valvular atrial fibrillation. Defendants' marketing campaign for Pradaxa® included promoting it as being more effective than warfarin in preventing stroke and systemic embolism, providing a convenient alternative to warfarin therapy because it does not require blood monitoring or dose adjustments, and does not require any dietary restrictions.
- 17. Defendants spent significant money in promoting Pradaxa®, which included \$67,000,000.00 spent during 2010 (although Pradaxa® was not approved for sale until October 19, 2010). ¹
- 18. During 2011, Defendants reportedly undertook 1.5 million Pradaxa® "detailing sessions" (marketing/sales visits by Defendants' sales force) with U.S. primary care physicians, internists, group practitioners, cardiologists, and practice nurses, spending approximately \$464,000,000.00 during this 12 month period to promote Pradaxa® in the United States.²
- 19. As part of their marketing of Pradaxa®, Defendants widely disseminated direct-to-consumer advertising campaigns that were designed to influence patients, including Ray Herndon Celsor, to make inquiries to their prescribing physician about Pradaxa® and/or request prescriptions for Pradaxa®.
- 20. In the course of these direct to consumer advertisements, Defendants overstated the efficacy of Pradaxa® with respect to preventing stroke and systemic

¹ Deborah Weinstein, Study: Sales Support is Dwindling, Not Dead, March 14, 2012, <u>Medical Marketing and Media</u>.

² Id.

embolism, failed to adequately disclose to patients that there is no drug, agent or means to reverse the anticoagulation effects of Pradaxa®, and that such irreversibility could have permanently disabling, life-threatening and fatal consequences.

- 21. Prior to Ray Herndon Celsor's prescription of Pradaxa®, Ray Herndon Celsor became aware of the promotional materials described herein.
- 22. Prior to Ray Herndon Celsor's prescription of Pradaxa®, Ray Herndon Celsor's prescribing physician received promotional materials and information from sales representatives of Defendants that Pradaxa® was more effective than warfain in reducing strokes in patients with non-valvular atrial fibrillation and was more convenient, without also adequately informing prescribing physicians that there was no reversal agent that could stop or control bleeding in patients taking Pradaxa®.
- 23. At all times relevant hereto, Defendants also failed to warn emergency room doctors, surgeons and other critical care medical professionals that unlike generally-known measures taken to treat and stabilize bleeding in users of warfarin, there is no effective agent to reverse the anticoagulation effects of Pradaxa®, and therefore no effective means to treat and stabilize patients who experience uncontrolled bleeding while taking Pradaxa®.
- 24. At all times relevant to this action, The Pradaxa® Medication Guide, prepared and distributed by Defendants and intended for U.S. patients to whom Pradaxa® has been prescribed, failed to warn and disclose to patients that there is no agent to reverse the anticoagulation effects of Pradaxa® and that if serious bleeding occurs, it may be irreversible, permanently disabling, and life-threatening.

- 25. From October 2010 until the end of March 2011, approximately 272,119 prescriptions for Pradaxa® were written in the United States. During that same period, there were 932 Pradaxa®-associated "Serious Adverse Event" ("SAE") Medwatch reports filed with the U.S. Food and Drug Administration, including at least 120 deaths and over 500 reports of severe, life-threatening bleeding.
- 26. From April 1 until the end of June 2011, there were an additional 856 Pradaxa®-associated "SAE" Medwatch reports filed with the U.S. Food and Drug Administration including at least 117 deaths and over 510 reports of severe, life-threatening bleeding.
- 27. During the Defendants' 2011 fiscal year, worldwide Pradaxa® sales eclipsed the \$1 billion threshold, achieving what is commonly known in the pharmaceutical industry as "blockbuster" sales status. ³
 - 28. Defendants' original labeling and prescribing information for Pradaxa®:
 - a. failed to disclose in the "Warnings" Section that there is no drug, agent or means to reverse the anticoagulation effects of Pradaxa®;
 - failed to advise prescribing physicians, such as Ray Herndon Celsor's physician, to instruct patients that there was no agent to reverse the anticoagulant effects of Pradaxa®;

8

³ Heide Oberhauser-Aslan and Tapan Sharma, Boehringer Sees Sales Rising Further as 2011 Profits Surge April 24, 2012 <u>WSJ.com</u>

- c. failed to investigate, research, study and consider, fully and adequately, patient weight as a variable factor in establishing recommended dosages of Pradaxa®;
- d. failed to investigate, research, study and define, fully and adequately,
 the safety profile of Pradaxa®;
- e. failed to provide adequate warnings about the true safety risks associated with the use of Pradaxa®;
- f. failed to warn that it is difficult or impossible to assess the degree and/or extent of anticoagulation in patients taking Pradaxa®;
- g. failed to provide adequate instructions on how to intervene and/or stabilize a patient who suffers a bleed while taking Pradaxa®;
- h. failed to provide adequate warnings regarding the need to assess renal functioning prior to starting a patient on Pradaxa® and to continue testing and monitoring of renal functioning periodically while the patient is on Pradaxa®;
- i. failed to provide adequate warnings and information related to the increased risks of bleeding events associated with aging patient populations of Pradaxa® users;
- j. failed to provide adequate warnings regarding the increased risk of gastrointestinal bleeds in those taking Pradaxa®, especially, in those patients with a prior history of gastrointestinal issues and/or upset;

- k. failed to include a "BOXED WARNING" about serious bleeding events associated with Pradaxa®;
- I. failed to include a "Bolded Warning" about serious bleeding events associated with Pradaxa®; and
- m. in their "Medication Guide" intended for distribution to patients to whom Pradaxa® has been prescribed, Defendants failed to disclose to patients that there is no drug, agent or means to reverse the anticoagulation effects of Pradaxa® and that if serious bleeding occurs, such irreversibility could have permanently disabling, life-threatening or fatal consequences.
- 27. During March, 2011, Defendants modified the U.S. labeling and prescribing information for Pradaxa®, which included additional information regarding the use of Pradaxa® in patients taking certain medications. Despite being aware of: (I) serious, and sometimes fatal, irreversible bleeding events associated with the use of Pradaxa®; (II) almost 1800 SAE Medwatch reports filed with the U.S. Food and Drug Administration, including at least 237 deaths and over 1,000 reports of severe, life-threatening bleeding, Defendants nonetheless failed to provide adequate disclosures or warnings in their label as detailed in Paragraphs 26 (a m).
- 28. On July 1, 2011, Pradaxa® was approved for sale in New Zealand with lower dosing (lowered from 150mg to 110mg twice a day) required for patients over 80 years of age and recommended for patients with moderate renal impairment.

- 29. On July 25, 2011, the <u>Archives of Internal Medicine</u> published The Use of Dabigatran [Pradaxa®] in Elderly Patients. [Vol 171, No. 14] which concluded that "The risk of major overdosage of...[Pradaxa®] in this [elderly] population is, however, much increased owing to frequent renal function impairment, low body weight, drug interactions that cannot be detected with a routine coagulation test and no antagonist available."
- 30. On January 21, 2011, Pradaxa® (under the brand name Prazaza®), in 75mg and 110mg doses only, is approved for sale in Japan to treat non-valvular atrial fibrillation.
- 31. On August 11, 2011, Japan's pharmaceutical regulatory authority announced that it was requiring a "BOXED WARNING" be added to Pradaxa® (marketed as Prazaza® in Japan) to call attention to reports of severe hemorrhages in patients treated with Pradaxa® (Prazaza®).
- 32. On September 1, 2011, the New Zealand pharmaceutical regulatory authority issued a "Prescriber Update" entitled "Dabigatran Is there a Bleeding Risk" in which physicians were alerted that Pradaxa® had a higher incidence of gastrointestinal bleeds than warfarin and that there was no reversal agent to neutralize the anticoagulation effects of Pradaxa®. A follow-up report issued in December 2011, indicated that among 10,000 New Zealanders who had taken Pradaxa®, there were 78 reports of serious bleeding events associated with Pradaxa® including 60 reports of gastrointestinal and rectal bleeding. Among the 78 serious events were 10 patient deaths and 55 hospitalizations. Three months later in March, 2012 the New England

Journal of Medicine published two letters from physicians in New Zealand addressing bleeding events associated with Pradaxa®. In one letter, physicians wrote, "We are concerned that the potential risks of this medication are not generally appreciated. The serious consequences of a lack of an effective reversal agent should not be underestimated."

- 33. During November 2011, Defendants modified the U.S. labeling and prescribing information for Pradaxa® adding additional information regarding the use of Pradaxa® in patients with kidney disease despite being aware of: (I) serious, and sometimes fatal, irreversible bleeding events associated with the use of Pradaxa®; (II) the July 25, 2011 article in the Archives of Internal Medicine; (III) the addition of a "BOXED WARNING" to Pradaxa® in Japan; and, (IV) the questions being raised by physicians in New Zealand about serious bleeding events associated with Pradaxa®, Defendants nonetheless failed to provide adequate disclosures or warnings in their label as detailed in Paragraphs 26 (a m).
- 34. On December 7, 2011, the U.S. Food and Drug Administration issued a Drug Safety Communication announcing that it was undertaking a "Drug Safety Review" of Post-Marketing Reports of Serious Bleeding Events with the anticoagulant Pradaxa. The purpose of the FDA's review is to determine if serious bleeding events associated with the use of Pradaxa® are more common than expected based on the Defendants' data submitted to the FDA.
- 35. As of December 31, 2011, the U.S. Food and Drug Administration received over 500 reports of deaths of people in the U.S. linked to Pradaxa® which, at

that point, had been available in the U.S. for approximately 14 months. In addition, there were over 900 reports of gastrointestinal hemorrhages, over 300 reports of rectal hemorrhages, and over 200 reports of cerebrovascular accidents suffered by U.S. citizens associated with Pradaxa®.

- 36. In January 2012, the Defendants modified the U.S. labeling and prescribing information for Pradaxa®. Despite being aware of: (i) serious, and sometimes fatal, irreversible bleeding events associated with the use of Pradaxa®; (ii) the July 25, 2011 article in the Archives of Internal Medicine; (iii) the addition of a "BOXED WARNING" to Pradaxa® in Japan; (iv) the questions being raised by physicians in New Zealand about serious bleeding events associated with Pradaxa®; and (v) the Drug Safety Communication published by the FDA in December, 2011, Defendants nonetheless failed to provide adequate disclosures or warnings in their label as detailed in Paragraphs 26 (a m).
- 37. During March 2012, in response to a directive from Health Canada, the governmental agency responsible for regulating pharmaceuticals in Canada, the Defendants' Canadian affiliate issued a "Dear Healthcare Provider" letter in which it advised Canadian healthcare providers of certain risks associated with the use of Pradaxa® (marketed as Pradax® in Canada) in elderly patients and patients with impaired kidney function and prosthetic heart valves. No such similar communication was sent to healthcare providers in the United States.
- 38. In April 2012, the Defendants modified the U.S. labeling and prescribing information for Pradaxa®. Despite being aware of: (i) serious, and sometimes fatal,

irreversible bleeding events associated with the use of Pradaxa®; (ii) the July 25, 2011 article in the Archives of Internal Medicine; (iii) the addition of a "BOXED WARNING" to Pradaxa® in Japan; (iv) the questions being raised by physicians in New Zealand about serious bleeding events associated with Pradaxa®; (v) the Drug Safety Communication published by the FDA in December, 2011; and (vi) the "Dear Healthcare Provider" letter Defendants were required to provide in Canada, Defendants nonetheless failed to provide adequate disclosures or warnings in their label as detailed in Paragraphs 26 (a – m).

39. At all times relevant hereto, Defendants failed to warn emergency room doctors, surgeons and other critical care medical professionals that unlike generally-known measures taken to treat and stabilize bleeding that occurs in the presence of warfarin, there is no effective agent to reverse the anticoagulation effects of Pradaxa® and therefore no effective means to treat and stabilize patients who experience uncontrolled bleeding while taking Pradaxa®.

Ray Herndon Celsor's use of Pradaxa® and resulting injuries

- 40. As a result of Defendants' claims regarding the effectiveness, safety, and benefits of Pradaxa®, Ray Herndon Celsor and Ray Herndon Celsor's physicians were unaware, and could not have reasonably known or have learned through reasonable diligence that Ray Herndon Celsor would be exposed to the risk of excessive and/or uncontrollable bleeding and the other risks and injuries described herein.
- 41. Therefore, Ray Herndon Celsor was prescribed Pradaxa® on or about March 19, 2011 for treatment of his medically necessary blood thinning needs. Shortly

thereafter, Ray Herndon Celsor suffered a severe gastrointestinal bleed on or about May 14, 2011 and underwent an esophagogastroduodenoscopy and colonoscopy on May 16, 2011 at his local hospital. After this procedure, Ray Herndon Celsor continued to decompensate and was transferred to Vanderbilt University Medical Center in Nashville, Tennessee and was admitted as a level 1 trauma patient. He was immediately taken to the operating room for exploratory laparotomy and esophagogastroduodenoscopy but he died during the procedure and did not respond to advanced cardiac life support protocol. Ray Herndon Celsor experienced excessive and/or uncontrollable bleeding, which was caused and/or worsened by Ray Herndon Celsor's use of Pradaxa®.

- 42. Prior to Ray Herndon Celsor's use of Pradaxa®, Defendants knew or should have known that the original labeling of the drug did not adequately warn Ray Herndon Celsor of the risks associated with using the drug as described above.
- 43. Prior to Ray Herndon Celsor's use of Pradaxa®, Defendants knew or should have known of the defective nature of Pradaxa® and persons who were prescribed and ingested Pradaxa® for even a brief period of time, including Ray Herndon Celsor, were at increased risk for developing life-threatening bleeds. Defendants, through their affirmative misrepresentations and omissions, concealed from Ray Herndon Celsor and Ray Herndon Celsor's physicians the true and significant risks associated with Pradaxa® use.
- 44. Ray Herndon Celsor was unaware of the increased risk for developing life-threatening injuries as compared to warfarin. Had Ray Herndon Celsor and/or Ray

Herndon Celsor's healthcare provider known of the risks and dangers associated with Pradaxa®, as well as the lack of additional benefits, and had Defendants provided adequate warnings that there is no agent to reverse the anticoagulation effects of Pradaxa®, Ray Herndon Celsor would not have used Pradaxa®.

45. As a direct and proximate result of using Pradaxa®, Ray Herndon Celsor has suffered severe personal injuries, physical pain and mental anguish, and ultimately, Ray Herndon Celsor died, all resulting from Ray Herndon Celsor's ingestion of Pradaxa®.

CAUSES OF ACTION

COUNT I STRICT LIABILITY-FAILURE TO WARN

- 46. Plaintiff hereby incorporates by reference all preceding paragraphs as if fully set forth herein.
- 47. At all times relevant to this suit, Defendants engaged in the business of designing, manufacturing, testing, marketing, labeling and placing into the stream of commerce Pradaxa® for sale to, and use by, members of the public.
- 48. At all times relevant to this suit, the dangerous propensities of Pradaxa® were known to Defendants, or were reasonably and scientifically knowable to them, through appropriate research and testing by known methods, at the time they distributed, supplied, or sold their respective product, and not known to ordinary physicians who would be expected to prescribe the drug for their patients.

- 49. The Pradaxa® manufactured by Defendants reached Plaintiff without substantial change and was ingested as directed.
- 50. Defendants marketed Pradaxa® in multiple ways, including but not limited to direct-to-consumer advertisements, which were misleading in that Defendants overstated the safety and efficacy of Pradaxa® and understated its risks.
- 51. The Pradaxa® was defective and unreasonably dangerous in that the labeling was insufficient to adequately warn physicians and users of the increased risk of excessive and/or uncontrollable bleeding.
- 52. As a direct and proximate result of the actions and inactions of the Defendants as set forth above, Plaintiff was exposed to Pradaxa® and suffered personal injuries, economic and non-economic damages including pain and suffering.
- 53. Defendants' actions and omissions as identified in this Complaint show that Defendants acted maliciously and/or intentionally disregarded Plaintiff's rights so as to warrant the imposition of punitive damages.

COUNT II STRICT PRODUCTS LIABILITY - DESIGN DEFECT

- 54. Plaintiff hereby incorporates by reference all preceding paragraphs as if fully set forth herein.
- 55. Defendants are the manufacturers, designers, distributers, sellers and suppliers of Pradaxa®, who sold Pradaxa® in the course of business.

- 56. The Pradaxa® manufactured, designed, sold, marketed, distributed, supplied and/or placed in the stream of commerce by Defendants was expected to and did reach the consumer without any alterations or changes.
- 57. The Pradaxa® administered to Plaintiff was defective in design or formulation in at least the following respects:
 - a. When it left the hands of the Defendants, this drug was unreasonably dangerous to an extent beyond that which could reasonably be contemplated by Plaintiff or Plaintiff's physicians;
 - b. Any benefit of this drug was outweighed by the serious and undisclosed risks of its use when prescribed and used as the Defendants intended;
 - c. The dosages and/or formulation of Pradaxa® sold by the Defendants was unreasonably dangerous;
 - d. There are no patients for whom the benefits of Pradaxa® outweighed the risks;
 - e. The product was not made in accordance with the Defendants' specifications or performance standards;
 - f. There are no patients for whom Pradaxa® is a safer and more efficacious drug than other drug products in its class; and/or
 - g. There were safer alternatives that did not carry the same risks and dangers that Defendants' Pradaxa® had.

- 58. The Pradaxa® administered to Plaintiff was defective at the time it was distributed by the Defendants or left their control.
- 59. The foreseeable risks associated with the design or formulation of the Pradaxa® include, but are not limited to, the fact that the design or formulation of Pradaxa® is more dangerous than a reasonably prudent consumer would expect when used in an intended or reasonably foreseeable manner, and/or did not have the claimed benefits.
- 60. The defective and unreasonably dangerous design and marketing of Pradaxa® was a direct, proximate and producing cause of Plaintiff's injuries and damages. Under strict products liability theories set forth in Restatement (Second) of Torts, Defendants are liable to Plaintiff for all damages claimed in this case.
- 61. As a direct, legal, proximate, and producing result of the defective and unreasonably dangerous condition of Pradaxa®, Plaintiff suffered personal injuries, economic and non-economic damages, including pain and suffering.
- 62. Defendants' actions and omissions as identified in this Complaint show that Defendants acted maliciously and/or intentionally disregarded Plaintiff's rights so as to warrant the imposition of punitive damages.

COUNT III NEGLIGENCE

63. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.

- 64. Defendants owed a duty to the general public and specifically to the Plaintiff to exercise reasonable care in the design, study, development, manufacture, promotion, sale, labeling, marketing and distribution of Pradaxa® at issue in this lawsuit.
- 65. Defendants breached their duty and failed to exercise reasonable care in the developing, testing, designing and manufacturing of Pradaxa® because, it was capable of causing serious personal injuries such as those suffered by Plaintiff during foreseeable use.
- 66. Defendants breached their duty and also failed to exercise reasonable care in the marketing of Pradaxa® because they failed to warn, that as designed, Pradaxa® was capable of causing serious personal injuries such as those suffered by Plaintiff during foreseeable use.
- 67. Defendants breached their duty and also failed to exercise ordinary care in the labeling of Pradaxa® and failed to issue to consumers and/or their health care providers adequate warnings of the risk of serious bodily injury or death due to the use of Pradaxa®. Moreover, Defendants over-promoted the benefits of Pradaxa® for anticoagulation therapy in patients suffering from atrial fibrillation and understated the risk of excessive and/or uncontrollable bleeding.
- 68. Defendants breached their duty and were negligent by, but not limited to, the following actions, misrepresentations, and omissions toward Plaintiff:

- a. In disseminating information to Plaintiff and Plaintiff's physicians that was negligently and materially inaccurate, misleading, false, and unreasonably dangerous to patients such as Plaintiff;
- b. Failing to conduct adequate pre-clinical and clinical testing and post-marketing surveillance to determine the safety of Pradaxa®;
- c. Failing to design and/or manufacture a product that could be used safely due to the lack of a known reversal agent; and
- d. In designing, manufacturing, and placing into the stream of commerce a product which was unreasonably dangerous for its reasonably foreseeable use, which Defendant knew or should have known could cause injury to Plaintiff.
- 69. Despite the fact that Defendants knew or should have known that Pradaxa® posed a serious risk of bodily harm to consumers and/or did not provide any additional benefits, Defendants continued to manufacture and market Pradaxa® for use by consumers.
- 70. Defendants knew or should have known that consumers, including Plaintiff, would foreseeably suffer injury as a result of Defendants' failure to exercise ordinary care as described above.
- 71. Defendants' failure to exercise reasonable care in the design, dosing information, marketing, warnings, labeling, and/or manufacturing of Pradaxa® was a proximate cause of Plaintiff's injuries and damages.

72. Defendants' conduct as described above, including but not limited to its failure to adequately test Pradaxa®, to provide adequate warnings, and its continued manufacture, sale and marketing of the product when it knew or should have known of the serious health risks it created, evidences actions and/or intentional disregard of the rights of Plaintiff so as to warrant the imposition of punitive damages.

COUNT IV NEGLIGENT MISREPRESENTATION AND/OR FRAUD

- 73. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.
- 74. Defendants represented that Pradaxa® was just as safe or safer and as effective or more effective than other anticoagulation alternatives and had additional benefits compared to other anticoagulation medications available on the market.
- 75. Defendants made these misrepresentations and actively concealed adverse information at a time when the Defendants knew, or should have known, that Pradaxa® had defects, dangers, and characteristics that were other than what Defendants had represented to Plaintiff and the health care industry generally. Specifically, Defendants misrepresented to and/or actively concealed from Plaintiff and the consuming public, among other things, that:
 - a. Pradaxa® had statistically significant increases in irreversible bleeds and other side effects which could result in serious, permanent injury or death;
 - b. Pradaxa® had not been fully or adequately tested;
 - c. Pradaxa® does not have any known reversal agents;

- d. Pradaxa® bleeds cannot be stopped or controlled by any effective medical processes or medical intervention;
- e. Failed to warn that it is difficult or impossible to assess the degree and/or extent of anticoagulation in patients taking Pradaxa®; and
- f. Pradaxa® was not as safe as blood thinners such as warfarin.
- 76. Defendants negligently and/or intentionally misrepresented or omitted this information in their product labeling, promotions and advertisements and instead labeled, promoted and advertised their product as safer and more effective than other types of anticoagulation alternatives and understated the risk of excessive and/or uncontrollable bleeding associated with Pradaxa®.
 - 77. The aforementioned misrepresentations were untrue and misleading.
- 78. Defendants knew or should have known that these representations were false and made the representations with the intent that Plaintiff and/or Plaintiff's prescribing physicians would rely on them, leading to the use of Pradaxa®.
- 79. At the time of Defendants' fraudulent misrepresentations, Plaintiff and/or Plaintiff's prescribing physicians were unaware of the falsity of the statements being made and believed them to be true. Plaintiff and/or Plaintiff's prescribing physicians justifiably relied on and/or were induced by the misrepresentations and/or active concealment and relied on the absence of safety information, which Defendants did suppress, conceal or failed to disclose, to Plaintiff's detriment.

- 80. As a direct and proximate result of the fraudulent acts and omissions, suppression and misrepresentation of Defendants, Plaintiff suffered personal injuries, economic and non-economic damages, including pain and suffering.
- 81. Defendants' actions and omissions as identified in this Complaint demonstrate malicious actions and/or intentional disregard of Plaintiff's rights so as to warrant the imposition of punitive damages.

COUNT V BREACH OF EXPRESS WARRANTY

- 82. Plaintiff incorporates by reference each preceding paragraph as though set forth fully at length herein.
- 83. Defendants expressly warranted, through their direct-to-consumer marketing, label, and sales representatives, that Pradaxa® was a safe and effective prescription blood thinner. The safety and efficacy of Pradaxa® constitute a material fact in connection with the marketing, promotion, and sale of Pradaxa®.
- 84. Pradaxa® manufactured and sold by Defendants did not conform to these express representations because it caused serious injury to consumers when taken in recommended dosages.
- 85. As a direct and proximate result of Defendants' breach of warranty, Plaintiff has suffered harm, damages and economic loss and will continue to suffer such harm, damages and economic loss in the future.

86. Defendants' actions and omissions as identified in this Complaint demonstrate malicious actions and/or intentional disregard of Plaintiff's rights so as to warrant the imposition of punitive damages.

COUNT VI BREACH OF IMPLIED WARRANTY

- 87. Plaintiff incorporates by reference each preceding paragraph as though set forth fully at length herein.
- 88. At the time Defendants researched, developed, designed, tested, manufactured, inspected, labeled, distributed, marketed, promoted, sold, and/or otherwise released Pradaxa® into the stream of commerce, Defendants knew of the use for which Pradaxa® was intended and impliedly warranted the product to be of merchantable quality and safe for such use.
- 89. Defendants breached their implied warranties of the Pradaxa® product sold to Plaintiff because this product was not fit for its common, ordinary, and intended use.
- 90. As a direct, foreseeable and proximate result of Defendants' breaches of implied warranties, Plaintiff suffered grievous bodily injury and consequential economic and other losses, as described above, when Plaintiff ingested Pradaxa®, in reasonable reliance upon the implied warranties.
- 91. Defendants' actions and omissions as identified in this Complaint demonstrate malicious actions and/or intentional disregard of Plaintiff's rights so as to warrant the imposition of punitive damages.

COUNT VII NEGLIGENCE PER <u>SE - DEFENDANTS' VIOLATION OF 21 U.S.C. §§ 331(a) & 352</u>

- 92. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.
- 93. As part of their duty to exercise reasonable care, Defendants were obligated to follow public laws and regulations enacted and promulgated to protect the safety of persons such as Plaintiff, including 21 U.S.C. §§ 331(a) & 352, and other statutes and regulations, which make it unlawful to misbrand prescription drug products.
- 94. The labeling, including package inserts, for Pradaxa® failed to conform to the requirements of 21 U.S.C. § 352, including subsections (a), (c), and (t), and the requirements of 21 C.F.R. § 201.100(c)(1), and, therefore, violated 21 U.S.C. § 331(a), which prohibits "[t]he introduction or delivery for introduction into interstate commerce of any food, drug, device, or cosmetic that is adulterated or misbranded."
- 95. Specifically, the product label and package insert for Pradaxa® is misbranded within the meaning of 21 U.S.C. § 352(a) and (f) because it was false and misleading and failed to give adequate warnings and directions for use by physicians who prescribe Pradaxa®.
- 96. Pradaxa® is misbranded pursuant to 21 U.S.C. § 352 because words, statements, or other information required by or under authority of chapter 21 U.S.C. § 352 are not prominently placed thereon with such conspicuousness and in such terms as

to render it likely to be read and understood by the ordinary individual under customary conditions of purchase and use.

- 97. Pradaxa® is misbranded pursuant to 21 U.S.C. § 352 because the labeling does not bear adequate directions for use, and/or the labeling does not bear adequate warnings against use where its use may be dangerous to health or against unsafe dosage or methods or duration of administration or application, in such manner and form as are necessary for the protection of users.
- 98. Pradaxa® is misbranded pursuant to 21 U.S.C. § 352 because it is dangerous to health when used in the dosage or manner, or with the frequency or duration prescribed, recommended, or suggested in the labeling thereof.
- 99. Because the Defendants each had a statutory duty under 21 U.S.C. § 352 (a) and (f) not to misbrand Pradaxa®, and because each of them violated this duty, they were guilty of negligence per se.
- 100. Pradaxa® is further misbranded pursuant to 21 C.F.R. § 201.56 because the labeling was not updated as new information became available that caused the labeling to become inaccurate, false, or misleading.
- 101. Defendants also violated 21 C.F.R. § 201.57 because they failed to identify specific tests needed for selection or monitoring of patients who took the prescription drug Pradaxa®.
- 102. Defendants violated 21 C.F.R. § 201.57 because the safety considerations regarding Pradaxa® are such that the drug should be reserved for certain situations, and the Defendants failed to state such information.

- 103. Pradaxa® is mislabeled pursuant to 21 C.F.R. § 201.57 because the labeling fails to describe serious adverse reactions and potential safety hazards, limitations in use imposed by it, and steps that should be taken if they occur.
- 104. Pradaxa® is mislabeled pursuant to 21 C.F.R. § 201.57 because the labeling was not revised to include a warning as soon as there was reasonable evidence of an association of a serious hazard with the drug (i.e., irreversible bleeding).
- 105. Pradaxa® is mislabeled pursuant to 21 C.F.R. § 201.57 because the labeling does not state an upper limit dosing beyond which safety and effectiveness have not been established.
- 106. Pradaxa® violates 21 C.F.R. § 210.122 because the labeling and packaging materials do not meet the appropriate specifications.
- 107. Pradaxa® violates 21 C.F.R. § 310.303 in that it is not safe and effective for its intended use.
- 108. Defendants violated 21 C.F.R. §§ 310.305 & 314.80 by failing to report adverse events associated with Pradaxa® as soon as possible or at least within 15 days of the initial receipt by the Defendants of the adverse drug experience.
- 109. Defendants violated 21 C.F.R. §§ 310.305 & 314.80 by failing to conduct an investigation of each adverse event associated with Pradaxa®, evaluate the cause of the adverse event, submit follow-up reports within the prescribed 15 calendar days of receipt of new information or as requested by the FDA, and keep records of the unsuccessful steps taken to seek additional information regarding serious, unexpected adverse drug experiences.

- 110. Defendants violated 21 C.F.R. § 314.80 by failing to provide periodic reports to the FDA containing (a) a narrative summary and analysis of the information in the report and an analysis of the IS-day Alert reports submitted during the reporting interval, (b) an Adverse Reaction Report for each adverse drug experience not already reported under the Post marketing IS-day Alert report, (c) a history of actions taken since the last report because of adverse drug experiences (for example, labeling changes or studies initiated) and/or (d) a copy of the published article from scientific or medical journals along with one or more I5-day Alert reports based on information from the scientific literature.
- 111. Defendants violated 21 C.F.R. § 312.32 because they failed to review all information relevant to the safety of Pradaxa® or otherwise received by Defendants from sources, foreign or domestic, including information derived from any clinical or epidemiological investigations, animal investigations, commercial marketing experience, reports in the scientific literature, and unpublished scientific papers, as well as reports from foreign regulatory authorities that have not already been previously reported to the agency by the sponsor.
- 112. Defendants failed to meet the standard of care set by the above statutes and regulations, which were intended for the benefit of individual consumers such as the Plaintiff, making Defendants liable to Plaintiff, and further, because each of them violated the above referenced duties required by these statutes and regulations, they are guilty of negligence per se.

113. Defendant's failure to adequately warn about the magnitude of the risk associated with use of Pradaxa® constitutes negligence per se. This negligence per se proximately caused injury to Plaintiff as described more fully herein.

COUNT VIII FRAUDULENT CONCEALMENT

- 114. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.
- 115. At all times during the course of dealings between Defendants and Plaintiff, and/or Plaintiff's healthcare providers, and/or the FDA, Defendants misrepresented the safety of Pradaxa® for its intended use.
- 116. Defendants knew or were reckless in not knowing that their representations were false.
- 117. In representations to Plaintiff, and/or Plaintiff's healthcare providers, and/or the FDA, Defendants fraudulently concealed and intentionally omitted the following material information:
 - a. that Pradaxa® was not as safe or effective as other forms of anticoagulation medication for atrial fibrillation patients;
 - that Defendants failed to investigate, research, study and consider, fully and adequately, patient weight as a variable factor in establishing recommended dosages of Pradaxa®;
 - c. that Defendants failed to investigate, research, study and define, fully and adequately, the safety profile of Pradaxa®;

- that Defendants failed to provide adequate warnings that there was no drug, agent or means to reverse the anticoagulation effects of Pradaxa®;
- e. that Defendants failed to include an adequate warning about serious bleeding events associated with Pradaxa®;
- f. that Defendants failed to warn it is difficult or impossible to assess the degree and/or extent of anticoagulation in patients taking Pradaxa®;
- g. that Defendants failed to adequately instruct physicians on how to intervene and/or stabilize a patient who suffers a bleed while taking Pradaxa®;
- h. that it is critical to fully assess renal functioning prior to starting a patient on Pradaxa® and to continue testing and monitoring of renal functioning periodically while the patient is on Pradaxa®;
- that there is an increased risk of bleeding events associated with aging patient populations of Pradaxa® users;
- j. that there is an increased risk of gastrointestinal bleeds in those taking Pradaxa®, especially, in those patients with a prior history of gastrointestinal issues and/or upset;
- k. that Pradaxa® was defective, and that it caused dangerous side effects, including but not limited to higher incidence of excessive and/or uncontrollable bleeding;
- I. that Pradaxa® was manufactured negligently;

- m. that Pradaxa® was manufactured defectively;
- n. that Pradaxa® was manufactured improperly;
- o. that Pradaxa® was designed negligently;
- p. that Pradaxa® was designed defectively; and
- g. that Pradaxa® was designed improperly.
- 118. Defendants were under a duty to disclose to Plaintiff, and Plaintiff's physicians, hospitals, healthcare providers, and/or the FDA the defective nature of Pradaxa®, including but not limited to the heightened risks of excessive and/or uncontrollable bleeding.
- 119. Defendants had sole access to material facts concerning the defective nature of the product and its propensity to cause serious and dangerous side effects, and hence, cause damage to persons who used Pradaxa®, including the Plaintiff, in particular.
- alia, the safety of Pradaxa® was made purposefully, willfully, wantonly, and/or recklessly, to mislead Plaintiff, and Plaintiff's physicians, hospitals and healthcare providers into reliance, continued use of Pradaxa®, and actions thereon, and to cause them to purchase, prescribe, and/or dispense Pradaxa® and/or use the product.
- 121. Defendants knew that Plaintiff and Plaintiff's physicians, hospitals, healthcare providers, and/or the FDA had no way to determine the truth behind Defendants' concealment and omissions, and that these included material omissions of facts surrounding Pradaxa®, as set forth herein.

- 122. Plaintiff, as well as Plaintiff's doctors, healthcare providers, and/or hospitals reasonably relied on facts revealed which negligently, fraudulently and/or purposefully did not include facts that were concealed and/or omitted by Defendants.
- 123. As a result of the foregoing acts and omissions the Plaintiff was and still is caused to suffer and/or is at a greatly increased risk of serious and dangerous side effects including, inter alia, excessive and/or uncontrollable bleeding, as well as other severe and personal injuries which are permanent and lasting in nature, physical pain and mental anguish, including diminished enjoyment of life.
- 124. As a result of the foregoing acts and omissions the Plaintiff requires and/or will require more health care and services and did incur medical, health, incidental and related expenses. Plaintiff further alleges that Plaintiff will in the future be required to obtain further medical and/or hospital care, attention, and services.
 - 125. By reason of the foregoing, Plaintiff has been damaged.

COUNT IX DECE<u>PTIVE TRADE PRACTICES</u>

- 126. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.
- 127. Defendants researched, developed, designed, tested, manufactured, inspected, labeled, distributed, marketed, promoted, sold, and/or otherwise released into the stream of commerce Pradaxa™, in the course of same, directly advertised or marketed the product to the FDA, health care professionals and consumers, including Plaintiff, or persons responsible for consumer.

- 128. Mr. Celsor purchased and used Pradaxa[™] for personal use and thereby suffered ascertainable losses as a result of Defendants' actions in violation of consumer protection laws, including the Tennessee Consumer Protection Act, Tenn. Code Ann. § 47-18-101 et. seq. which include the following violations:
- a. Representing that goods or services have characteristics, ingredients, uses, benefits or qualities that they do not have;
- b. Advertising goods or services with the intent not to sell them as advertised;
- c. Representing that goods have sponsorship, approval, characteristics, ingredients, uses, benefits or quantities which they do not have;
- d. Failing to disclose information concerning goods which was known at the time of the transaction if such failure to disclose such information was intended to induce the consumer into a transaction into which the consumer would not have entered had the information been disclosed;
 - e. Unconscionable actions and courses of action; and
- f. Engaging in fraudulent or deceptive conduct that creates a likelihood of confusion or misunderstanding.
- 129. Defendants uniformly communicated the purported benefits of Pradaxa[™] while failing to disclose the serious and dangerous side-effects related to Pradaxa[™] and of the true state of Pradaxa[™], the regulatory status, its safety, its efficacy and its true usefulness. Defendants made these representations to physicians, the medical

community at large and to patients and consumers, such as Plaintiff, in their marketing and advertising.

- 130. Defendants' conduct in connection with Pradaxa[™] was also impermissible and illegal in that it created a likelihood of confusion and misunderstanding because Defendants misleadingly, falsely and/or deceptively misrepresented and omitted numerous material facts regarding the utility, benefits, costs, safety, efficacy and advantages of Pradaxa[™].
- 131. As a direct, proximate and foreseeable result of Defendants' statutory violations, Plaintiff suffered the injuries and consequential economic and other losses, as described above, when Plaintiff ingested Pradaxa™.

COUNT X LOSS OF CONSORTIUM

- 132. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.
- 133. Plaintiff, Amanda Scott, was at all times relevant hereto the daughter of Plaintiff.
- 134. For the reasons set forth herein, Plaintiff has been caused, presently and in the future, to suffer the loss of her father's companionship and society, and accordingly, the Plaintiff has been caused great mental anguish.

PUNITIVE DAMAGES

135. Plaintiff hereby incorporates by reference all of the above allegations as if fully set forth herein.

- 136. At all material times, the Defendants knew or should have known that Pradaxa® was inherently dangerous.
- 137. Despite their knowledge, the Defendants continued to aggressively market Pradaxa® to consumers, including Plaintiff, without disclosing its dangerous side effects when there existed safer alternative products.
- 138. Despite Defendants' knowledge of Pradaxa®'s defective and unreasonably dangerous nature, Defendants continued to test, design, develop, manufacture, label, package, promote, market, sell and distribute it so as to maximize sales and profits at the expense of the health and safety of the public, including the Plaintiff, in conscious disregard of the foreseeable harm caused by Pradaxa®.
 - 139. Defendants' conduct was intentional and/or wanton.
- 140. Defendants' conduct as described above, including, but not limited to, their failure to adequately test their product, to provide adequate warnings, and their continued manufacture, sale, and marketing or their products when they knew or should have known of the serious health risks created, evidences a flagrant disregard of human life as to warrant the imposition of punitive damages as the acts or omissions were committed with knowing, conscious and deliberate disregard for the rights and safety of consumers, including Plaintiff.

PRAYER FOR RELIEF

WHEREFORE, Plaintiff prays for relief as follows:

 Actual damages as alleged, jointly and/or severally against Defendants, in excess of \$75,000.00;

- 2. Medical expenses and other economic damages in an amount to be determined at trial of this action;
- 3. Pain and suffering;
- 4. Wrongful death;
- 5. Burial and funeral expenses;
- 6. Loss of companionship and society;
- 7. Punitive damages alleged against Defendants, including Plaintiff's attorney fees, in excess of \$75,000.00;
- 8. Prejudgment interest at the highest lawful rate allowed by law;
- Interest on the judgment at the highest legal rate from the date of judgment until collected;
- 10. Attorneys' fees, expenses, and costs of this action; and
- 11. Such further relief as this Court deems necessary, just and proper.

JURY DEMAND

Plaintiff hereby demands a trial by jury on all issues so triable.

Dated this 14th day of May, 2012

Respectfully submitted,

/s/ Lee L. Coleman

Lee L. Coleman

Lee L. Coleman
TN Bar No. 019689
HUGHES & COLEMAN
P.O. Box 10120
Bowling Green, KY 42102

Telephone: (270) 785-2110

Facsimile: (270-782-8820)

WATTS GUERRA CRAFT LLP Mikal C. Watts (pro hac motion pending) Federal Bar No. 12419 Ryan L. Thompson (pro hac motion pending) Federal Bar No. 602642 2506 North Port Avenue Corpus Christi, Texas 78401 Telephone: 361.693.3100

Fax: 361.882.1261

Email: mcwatts@wgclawfirm.com Email: rthompson@wgclawfirm.com

ATTORNEYS FOR PLAINTIFFS

CIVIL COVER SHEET

The JS 44 civil cover sheet and the information contained herein neither replace nor supplement the filing and service of pleadings or other papers as required by law, except as provided by local rules of court. This form, approved by the Judicial Conference of the United States in September 1974, is required for the use of the Clerk of Court for the purpose of initiating the civil docket sheet. (SEE INSTRUCTIONS ON THE REVERSE OF THE FORM.)

the civil docket sheet. (SEE h	birderions on the Revense of The Forest,				
I. (a) PLAINTIFFS			DEFENDANTS		
	ndividually and as Representative of t erndon Celsor, Deceased	he =	Boehringer Ingel	heim Pharmaceuticals	s, Inc. et al
-	of First Listed Plaintiff Montgomery		County of Residence of	That Listed Detendant	airfield
	(CEPT IN U.S. PLAINTIFF CASES)			(IN U.S. PLAINTIFF CASES O	
				CONDEMNATION CASES, USI IVOLVED.	E THE LOCATION OF THIS
• •	Address, and Telephone Number)		Attorneys (If Known)		
Lee Coleman, H Bowling Green,	ughes & Coleman, P.O. Box 10120,	+	i		
II. BASIS OF JURISD		III. CI	TIZENSHIP OF PI	RINCIPAL PARTIES	Place an "X" in One Box for Plaintiff
☐ 1 U.S. Government	□ 3 Federal Question	((For Diversity Cases Only) PT		and One Box for Defendant) PTF DEF
Plaintiff	(U.S. Government Not a Party)	Citize	en of This State	1	
2 U.S. Government	📜 4 Diversity	Citize	en of Another State	2	
Defendant	(Indicate Citizenship of Parties in Item III)				1 6 1 6
			en or Subject of a reign Country	3 🗖 3 Foreign Nation	
IV. NATURE OF SUI	T (Place an "X" in One Box Only)	i ior	Descriptions of the second		STEER TO BE STEER TO
☐ 110 Insurance	TORTS PERSONAL INJURY PERSONAL INJURY		0 Agriculture	☐ 422 Appeal 28 USC 158	☐ 400 State Reapportionment
120 Marine	☐ 310 Airplane ☐ 362 Personal Injury	- 🗇 62	20 Other Food & Drug	☐ 423 Withdrawal 28 USC 157	☐ 410 Antitrust ☐ 430 Banks and Banking
130 Miller Act	☐ 315 Airplane Product Med. Malpractic Liability 365 Personal Injury		25 Drug Related Seizure of Property 21 USC 881	26 USC 137	450 Commerce
☐ 140 Negotiable Instrument☐ 150 Recovery of Overpayment	☐ 320 Assault, Libel & Product Liabilit	ty 🗇 63	30 Liquor Laws	PROPERTYRICHUS	☐ 460 Deportation ☐ 470 Racketeer Influenced and
& Enforcement of Judgmen	Slander 368 Asbestos Person		10 R.R. & Truck	820 Copyrights 830 Patent	Corrupt Organizations
☐ 151 Medicare Act ☐ 152 Recovery of Defaulted	330 Federal Employers' Injury Product Liability Liability		50 Airline Regs. 50 Occupational	☐ 840 Trademark	☐ 480 Consumer Credit
Student Loans	☐ 340 Marine PERSONAL PROPE	RTY	Safety/Health		490 Cable/Sat TV 810 Selective Service
(Excl. Veterans)	☐ 345 Marine Product ☐ 370 Other Fraud Liability ☐ 371 Truth in Lendin		90 Other		☐ 850 Securities/Commodities/
153 Recovery of Overpayment of Veteran's Benefits	Liability 371 Truth in Lendin 350 Motor Vehicle 380 Other Personal		10 Fair Labor Standards	☐ 861 HIA (1395ff)	Exchange
☐ 160 Stockholders' Suits	355 Motor Vehicle Property Damag		Act 20 Labor/Mgmt. Relations	862 Black Lung (923) 863 DIWC/DIWW (405(g))	875 Customer Challenge 12 USC 3410
☐ 190 Other Contract ☐ 195 Contract Product Liability	Product Liability 385 Property Damag	· .	30 Labor/Mgmt.Reporting	☐ 864 SSID Title XVI	☐ 890 Other Statutory Actions
☐ 196 Franchise	Injury		& Disclosure Act	☐ 865 RSI (405(g)) FEDERAL TAX SUEES	891 Agricultural Acts 892 Economic Stabilization Act
REAL PROPERTY.	☐ 441 Voting ☐ 510 Motions to Vac		40 Railway Labor Act 90 Other Labor Litigation	☐ 870 Taxes (U.S. Plaintiff	☐ 893 Environmental Matters
☐ 210 Land Condemnation ☐ 220 Foreclosure	☐ 441 Voting ☐ 510 Motions to Vac ☐ 442 Employment Sentence	0 7	91 Empl. Ret. Inc.	or Defendant)	894 Energy Allocation Act
230 Rent Lease & Ejectment	☐ 443 Housing/ Habeas Corpus:	ļ	Security Act	☐ 871 IRS—Third Party 26 USC 7609	☐ 895 Freedom of Information Act
240 Torts to Land	Accommodations 530 General 535 Death Penalty		IMMIGRATION	20 030 7007	900Appeal of Fee Determination
245 Tort Product Liability290 All Other Real Property	445 Amer. w/Disabilities - 540 Mandamus & C		62 Naturalization Application		Under Equal Access to Justice
2 2,0114 0 444 7	Employment		.63 Habeas Corpus - Alien Detainee		950 Constitutionality of
	☐ 446 Amer. w/Disabilities - ☐ 555 Prison Condition	on 4	65 Other Immigration		State Statutes
	☐ 440 Other Civil Rights		Actions		
▼1 Original □ 2	e an "X" in One Box Only) Removed from	Rec	nstated or 3 anoth	er district	n Judgment
Troccouning	Cite the U.S. Civil Statute under which you 28 US Section 1332	are filing	(Do not cite jurisdiction	al statutes unless diversity):	
VI. CAUSE OF ACT	Brief description of cause: Wrongful death; product liability				
VII. REQUESTED I COMPLAINT:			DEMAND \$	CHECK YES only JURY DEMAND	y if demanded in complaint: e: ☑ Yes ☐ No
VIII. RELATED CA	SE(S) (See instructions): JUDGE			DOCKET NUMBER	
	SIGNATURE OF	ATTORNE	Y OF RECORD		
DATE 05/14/2012	/s/ Lee Colen				
FOR OFFICE USE ONLY	.5. 250 0000				
RECEIPT#	AMOUNT APPLYING IFF	P	JUDGE	мад. л	UDGE

for the

	Wildele Disc	inct of Tellifessee	
Amanda Scott, Individually and the Estate of Ray Herndon)	
Plaintiff)	
v.) Civil Action No.	
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)	
Defendant		,	
	SUMMONS IN	A CIVIL ACTION	
To: (Defendant's name and address)	Boehringer Ingelheim Pha c/o CT Corporation Syster 800 S. Gay Street, Ste. 20 Knoxville, Tennessee 379	n 121	
are the United States or a Unite P. 12 (a)(2) or (3) — you must	rvice of this summons on y d States agency, or an offic serve on the plaintiff an ans	ou (not counting the day you receiver or employee of the United States on the attached complaint or must be served on the plaintiff	tes described in Fed. R. Civ. a motion under Rule 12 of
If you fail to respond, j You also must file your answer		entered against you for the relief	demanded in the complaint.
		CLERK OF COURT	
Date:		Signature of	Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

	This summons for (nan	ne of individual and title, if any)			
was re	ceived by me on (date)	•			-
	☐ I personally served	the summons on the individual at (place)		
			on (date)	; or	
	☐ I left the summons	at the individual's residence or usu	al place of abode with (name)		
		, a person of	suitable age and discretion who resid	des there,	
	on (date)	, and mailed a copy to the	individual's last known address; or		
	☐ I served the summo	ons on (name of individual)		,,	who is
	designated by law to a	accept service of process on behalf	of (name of organization)		
			on (date)	; or	
	☐ I returned the summ	nons linevectited because			; or
	Other (specify):				
	My fees are \$	for travel and \$	for services, for a total of \$	0.00	·
	I declare under penalt	y of perjury that this information is	true.		
Date:			Server's signature		
			U		
		4	Printed name and title		
			Server's address		***************************************

for the

Amanda Scott, Individually and)
the Estate of Ray Herndon Plaintiff	Celsor, Deceased	– <u> </u>
v.) Civil Action No.
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)
	Province Asia Salahan Salah	-) -)
Defendant		,
	SUMMONS	S IN A CIVIL ACTION
To: (Defendant's name and address)	Boehringer Ingelheim Co CT Corporation Sy 800 S. Gay Street, Ste Knoxville, Tennessee	ystem e. 2021
are the United States or a Unite P. 12 (a)(2) or (3) — you must	rvice of this summons of d States agency, or an of serve on the plaintiff ar	
If you fail to respond, j You also must file your answer	udgment by default wil	Ill be entered against you for the relief demanded in the complaint.
Date:		
		Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

	This summons for (no	nme of individual and title, if	any)			
was rec	ceived by me on (date)		·			
	☐ I personally served	d the summons on the in	ndividual at (place)			
				on (date)	; or	
	☐ I left the summons	s at the individual's resi	idence or usual pla	ce of abode with (name)		
		,	, a person of suitab	le age and discretion who resid	les there,	
	on (date)	, and mailed	a copy to the indiv	ridual's last known address; or		
	☐ I served the summ	ons on (name of individual	<i>I</i>)		, w	ho is
	designated by law to	accept service of proce				
				on (date)	; or	
	☐ I returned the sum	mons unexecuted becau	use			; or
	☐ Other (specify):					
	My fees are \$	for travel and	d\$	for services, for a total of \$	0.00	
	I declare under penal	ty of perjury that this ir	nformation is true.			
Date:						
Date.		_		Server's signature		
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for the

	Wildelf Die	
Amanda Scott, Individually an the Estate of Ray Herndor)
Plaintiff	•)
v.) Civil Action No.
Boehringer Ingelheim Pharn	naceuticals, Inc., et al)
Defendan	it)
	SUMMONS IN	A CIVIL ACTION
To: (Defendant's name and address)	Boehringer Ingelheim Cor c/o CT Corporation Syster One Corporate Center Hartford, CT 06103	
are the United States or a United P. 12 (a)(2) or (3) — you must	ervice of this summons on yed States agency, or an office serve on the plaintiff an an	rou (not counting the day you received it) — or 60 days if you per or employee of the United States described in Fed. R. Civ. swer to the attached complaint or a motion under Rule 12 of on must be served on the plaintiff or plaintiff's attorney,
If you fail to respond, You also must file your answe		entered against you for the relief demanded in the complaint.
		CLERK OF COURT
Date:		
	•	Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

		(name of individual and title, ij	f any)			
was rec	ceived by me on (date	2)	*			
	☐ I personally serv	ved the summons on the i	individual at (place)			
				on (date)	; or	
	☐ I left the summo	ons at the individual's res	idence or usual pla	ce of abode with (name)		
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	on (date)	, and mailed	a copy to the indiv	vidual's last known address; or		
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	☐ I returned the su	ımmons unexecuted beca	use		; 0:	r
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	My fees are \$	for travel an	nd \$	for services, for a total of \$	0.00	_
	I declare under per	nalty of perjury that this is	nformation is true.			
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		-		Server's address		-

for the

	Wilddie D	district of Telmessee
Amanda Scott, Individually and the Estate of Ray Herndon)
Plaintiff)
v.) Civil Action No.
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)
Defendant)
	SUMMONS I	IN A CIVIL ACTION
To: (Defendant's name and address)	Boehringer Ingelheim US c/o CT Corporation Syst One Corporate Center Hartford, CT 06103	
are the United States or a Unite P. 12 (a)(2) or (3) — you must	rvice of this summons on d States agency, or an off serve on the plaintiff an a	
If you fail to respond, j You also must file your answer		be entered against you for the relief demanded in the complaint.
		CLERK OF COURT
Date:		
		Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

was re	ceived by me on (date)	•		
	☐ I personally served	the summons on the individual at (place)		
			on (date)	; or
	☐ I left the summons	at the individual's residence or usual pla	ace of abode with (name)	
		, a person of suitab	ole age and discretion who resid	les there,
	on (date)	, and mailed a copy to the indi-	vidual's last known address; or	
	☐ I served the summe	ons on (name of individual)		, who is
	designated by law to	accept service of process on behalf of (no		
			on (date)	; or
	☐ I returned the sum	mons unexecuted because		; or
	Other (specify):			
	My fees are \$	for travel and \$	for services, for a total of \$	0.00
	I declare under penalt	y of perjury that this information is true.		
Date:			Server's signature	
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for the

	Wilduic D	istrict of Tellifessee
Amanda Scott, Individually and the Estate of Ray Herndon)
Plaintiff)
v.) Civil Action No.
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)
Defendant)
	SUMMONS	N A CIVIL ACTION
To: (Defendant's name and address)	Boehringer Ingelheim P Binger Strasse 173	narma GmbH & Co. KG
	55216 Ingelheim am Rh	ein, Germany
are the United States or a Unite P. 12 (a)(2) or (3) — you must	ervice of this summons or d States agency, or an of serve on the plaintiff an	a you (not counting the day you received it) — or 60 days if you ficer or employee of the United States described in Fed. R. Civanswer to the attached complaint or a motion under Rule 12 of action must be served on the plaintiff or plaintiff's attorney,
If you fail to respond, j You also must file your answer	udgment by default will or motion with the cour	be entered against you for the relief demanded in the complaint
		CLERK OF COURT
Dota		
Date:		Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

	This summons for (nan	ne of individual and title, if any)		
was re	ceived by me on (date)			
	☐ I personally served	the summons on the individual at (place)	
			on (date)	
	☐ I left the summons	at the individual's residence or usu	al place of abode with (name)	
		, a person of s	suitable age and discretion who resid	les there,
	on (date)	, and mailed a copy to the	individual's last known address; or	
	☐ I served the summo	ons on (name of individual)		, who is
	designated by law to a	accept service of process on behalf		
			on (date)	; or
	☐ I returned the summ	nons unexecuted because		; or
	Other (specify):			
	My fees are \$	for travel and \$	for services, for a total of \$	0.00
	I declare under penalty	y of perjury that this information is	true.	
Date:			Server's signature	
			5	
			Printed name and title	
			Server's address	

for the

Amanda Scott, Individually and the Estate of Ray Herndon		
Plaintiff)	
v.)	Civil Action No.
Boehringer Ingelheim Pharm	aceuticals, Inc., et al	
Defendant)	
	SUMMONS IN A CI	VIL ACTION
To: (Defendant's name and address)	Boehringer Ingelheim Internation Binger Strasse 173 55216 Ingelheim am Rhein, Ger	
are the United States or a Unite P. 12 (a)(2) or (3) — you must	rvice of this summons on you (no d States agency, or an officer or e serve on the plaintiff an answer to	ot counting the day you received it) — or 60 days if you employee of the United States described in Fed. R. Civ. to the attached complaint or a motion under Rule 12 of st be served on the plaintiff or plaintiff's attorney,
If you fail to respond, j You also must file your answer		ed against you for the relief demanded in the complaint.
		CLERK OF COURT
Date:		
		Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

	This summons for (nan	ne of individual and title, if any)			
was re	ceived by me on (date)	•			
	☐ I personally served	the summons on the individual at (1	place)		
		W	on (date)	; or	
	☐ I left the summons	at the individual's residence or usua	al place of abode with (name)		
		, a person of s	suitable age and discretion who resid	des there,	
	on (date)	, and mailed a copy to the	individual's last known address; or		
	☐ I served the summo	ons on (name of individual)		,,	who is
	designated by law to a	accept service of process on behalf		***	
			on (date)	; or	
	☐ I returned the summ	nons unexecuted because			; or
	☐ Other (specify):				
	My fees are \$	for travel and \$	for services, for a total of \$	0.00	·
	I declare under penalty	y of perjury that this information is	true.		
Date:					
			Server's signature		
			Printed name and title		
			Server's address		

or the

Middle I	District of Tennessee
Amanda Scott, Individually and as Representative of the Estate of Ray Herndon Celsor, Deceased Plaintiff v. Boehringer Ingelheim Pharmaceuticals, Inc., et al Defendant)) Civil Action No.))
SUMMONS	IN A CIVIL ACTION
To: (Defendant's name and address) Boehringer Ingelheim F Binger Strasse 173 55216 Ingelheim am Ri	
are the United States or a United States agency, or an o P. 12 (a)(2) or (3) — you must serve on the plaintiff an	
If you fail to respond, judgment by default will You also must file your answer or motion with the cou	! be entered against you for the relief demanded in the complaint. art.
Date: MAY 1 4 2012	CLERK OF COURT Signature of Clerk or Deputy Clerk

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

This summons for (no	ame of individual and title, if any)		
eceived by me on (date)			
☐ I personally serve	d the summons on the individual a	t (place)	
		on (date)	; or
☐ I left the summon	s at the individual's residence or us	sual place of abode with (name)	
	, a person o	of suitable age and discretion who resid	les there,
on (date)	, and mailed a copy to t	he individual's last known address; or	
☐ I served the sumn	nons on (name of individual)		, who
designated by law to	accept service of process on beha		
		on (date)	; or
☐ I returned the sun	nmons unexecuted because		; (
☐ Other (specify):			
in Callet (opensy).			
My fees are \$	for travel and \$	for services, for a total of \$	0.00
I declare under pena	lty of perjury that this information	is true.	
		Server's signature	
		Printed name and title	
		2 7 111000 11111111111111111111111111111	
		Server's address	
		SCI FOI B GGGI USS	

for the

W div				
1	Middle District of Tennessee			
Amanda Scott, Individually and as Representa the Estate of Ray Herndon Celsor, Deceas				
Plaintiff)			
v.) Civil Action No.			
Boehringer Ingelheim Pharmaceuticals, Inc.,	et al)			
Defendant)			
SUM	MONS IN A CIVIL ACTION			
Binger Strasse	elheim International GmbH 173 m am Rhein, Germany			
A lawsuit has been filed against you. Within 21 days after service of this summons on you (not counting the day you received it) — or 60 days if you are the United States or a United States agency, or an officer or employee of the United States described in Fed. R. Civ. P. 12 (a)(2) or (3) — you must serve on the plaintiff an answer to the attached complaint or a motion under Rule 12 of the Federal Rules of Civil Procedure. The answer or motion must be served on the plaintiff or plaintiff's attorney, whose name and address are: Lee L. Coleman HUGHES & COLEMAN P.O. Box 10120 Bowling Green, KY 42102				
If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.				
Date: MAY 1 4 2012	CLERK OF COURT Signature of Clerk or Deputy Clerk			

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

was re	This summons for (nan	me of individual and title, if any)			
	☐ I personally served	i the summons on the individual at (place)		
			on (date)	; or	
	☐ I left the summons	at the individual's residence or usual pl			
			ble age and discretion who resid		
	on (date), and mailed a copy to the individual's last known address; or				
	☐ I served the summe	ons on (name of individual)		, who	is
	designated by law to	accept service of process on behalf of (n	ame of organization)		
			on (date)	; or	
	☐ I returned the summ	mons unexecuted because		; ·	or
	Other (specify):				
	My fees are \$	for travel and \$	for services, for a total of \$	0.00	
	I declare under penalt	y of perjury that this information is true			
Date:					
			Server's signature		_
			Printed name and title		-
			Server's address		

for the

for the				
	Middle District of Tennessee			
Amanda Scott, Individually and as Representa the Estate of Ray Herndon Celsor, Deceas				
Plaintiff)			
v.) Civil Action No.			
Boehringer Ingelheim Pharmaceuticals, Inc.,	et al			
Defendant				
SUM	MONS IN A CIVIL ACTION			
c/o CT Corpora 800 S. Gay Stre	To: (Defendant's name and address) Boehringer Ingelheim Vetmedica, Inc. c/o CT Corporation System 800 S. Gay Street, Ste. 2021 Knoxville, Tennessee 37929-9710			
A lawsuit has been filed against you.				
Within 21 days after service of this summons on you (not counting the day you received it) — or 60 days if you are the United States or a United States agency, or an officer or employee of the United States described in Fed. R. Civ. P. 12 (a)(2) or (3) — you must serve on the plaintiff an answer to the attached complaint or a motion under Rule 12 of the Federal Rules of Civil Procedure. The answer or motion must be served on the plaintiff or plaintiff's attorney, whose name and address are: Lee L. Coleman HUGHES & COLEMAN P.O. Box 10120 Bowling Green, KY 42102				
If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.				
	CLERK OF COURT			
Date: MAY 1 4 2012	Ple			
	Signature of Clerk or Deputy Clerk			

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

was re	This summons for (nanceived by me on (date)	ne of individual and title, if any)			
	☐ I personally served	the summons on the individual at (place)			
			on (date)	; or -	
	☐ I left the summons	at the individual's residence or usual pla	ace of abode with (name)		
		, a person of suital	ble age and discretion who resid	des ther	e,
	on (date)	, and mailed a copy to the indi	vidual's last known address; or		
		ONS ON (name of individual)			, who is
	designated by law to a	accept service of process on behalf of (no	ume of organization)		
			On (date)	; or	
	☐ I returned the summ	nons unexecuted because			; or
	Other (specify):				
	My fees are \$	for travel and \$	for services, for a total of \$		0.00
	I declare under penalty	y of perjury that this information is true.			
Date:					
			Server's signature		
		· · · · · · · · · · · · · · · · · · ·	Printed name and title		
			Server's address		<u> </u>

UNITED STATES DISTRICT COURT for the

		101 tile		
	Middle l	District of Te	ennessee	
Amanda Scott, Individually and the Estate of Ray Herndon)		
Plaintiff)		
v.) c	Civil Action No.	
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)		
Defendant		, j		
	SUMMONS	IN A CIVIL	L ACTION	
To: (Defendant's name and address)	Boehringer Ingelheim F c/o CT Corporation Sys 800 S. Gay Street, Ste Knoxville, Tennessee S	stem . 2021	cals, Inc.	
A lawsuit has been filed against you. Within 21 days after service of this summons on you (not counting the day you received it) — or 60 days if you are the United States or a United States agency, or an officer or employee of the United States described in Fed. R. Civ. P. 12 (a)(2) or (3) — you must serve on the plaintiff an answer to the attached complaint or a motion under Rule 12 of the Federal Rules of Civil Procedure. The answer or motion must be served on the plaintiff or plaintiff's attorney, whose name and address are: Lee L. Coleman HUGHES & COLEMAN P.O. Box 10120 Bowling Green, KY 42102				
If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.				
Date: MAY 1 4 201	2		CLERK OF COURT Signature of Clerk or Deputy Clerk	

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

	This summons for (na	me of individual and title, if a	any)			
was re	ceived by me on (date)					
	☐ I personally served	the summons on the inc	dividual at (place)	<u></u>		
			on (date)	; or		
	☐ I left the summons at the individual's residence or usual place of abode with (name)					
		, , ,	a person of suitable age and discretion who resid	les there	,	
	on (date)	, and mailed a	copy to the individual's last known address; or			
	☐ I served the summ	ons on (name of individual)			, who is	
	designated by law to	accept service of process	s on behalf of (name of organization)		-	
			on (date)	; or		
	☐ I returned the sum	mons unexecuted becaus	se		; or	
	Other (specify):			1		
	My fees are \$	for travel and	\$ for services, for a total of \$	0	.00	
	I declare under penalt	ty of perjury that this inf	formation is true.			
Datas						
Date:			Server's signature	,		
		_	Printed name and title			
		_	Server's address			

for the

		101 410		
	Middle D	District of Te	nnessee	
Amanda Scott, Individually and the Estate of Ray Herndon	l as Representative of Celsor, Deceased)		
Plaintiff)		
v.) Ci	ivil Action No.	
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)		
Defendant)		
	SUMMONS 1	IN A CIVIL	ACTION	
To: (Defendant's name and address)	Boehringer Ingelheim U c/o CT Corporation Sys One Corporate Center Hartford, CT 06103		tion	
A lawsuit has been filed against you. Within 21 days after service of this summons on you (not counting the day you received it) — or 60 days if you are the United States or a United States agency, or an officer or employee of the United States described in Fed. R. Civ. P. 12 (a)(2) or (3) — you must serve on the plaintiff an answer to the attached complaint or a motion under Rule 12 of the Federal Rules of Civil Procedure. The answer or motion must be served on the plaintiff or plaintiff's attorney, whose name and address are: Lee L. Coleman HUGHES & COLEMAN P.O. Box 10120				
Bowling Green, KY 42102 If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.				
MAY 1 4 2	20 12 		CLERK OF COURT Signature of Clerk or Deputy Clerk	

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

vas rec	ceived by me on (date)					
	☐ I personally served	the summons on the individual at (place)				
	W		on (date)	_ ; or		
	☐ I left the summons at the individual's residence or usual place of abode with (name)					
		, a person of suitab	ole age and discretion who resid	les there,		
	on (date)	, and mailed a copy to the indi-	vidual's last known address; or			
	☐ I served the summo	ns on (name of individual)		, who	is	
	designated by law to a	accept service of process on behalf of (na				
			on (date)	_ ; or		
	☐ I returned the summ	nons unexecuted because		;	or	
	☐ Other (specify):					
	My fees are \$	for travel and \$	for services, for a total of \$	0.00	·	
	I declare under penalty	y of perjury that this information is true.				
Date:			Server's signature			
			Printed name and title	-	_	
			Server's address	· · · · · · · · · · · · · · · · · · ·	_	

for the					
	Middle I	District of T	ennessee		
Amanda Scott, Individually an the Estate of Ray Herndon)			
Plaintiff)			
v.) (Civil Action No.		
Boehringer Ingelheim Pharm	aceuticals, Inc., et al)			
Defendani)			
	SUMMONS I	IN A CIVI	L ACTION		
To: (Defendant's name and address)	Boehringer Ingelheim C c/o CT Corporation Sys One Corporate Center Hartford, CT 06103	orporation tem			
A lawsuit has been file Within 21 days after se	rvice of this summons or	n you (not c	counting the day you received it) — or 60 days if you		
P. 12 (a)(2) or (3) — you must	serve on the plaintiff an	answer to the training to the contract to the	ployee of the United States described in Fed. R. Civ. he attached complaint or a motion under Rule 12 of the served on the plaintiff or plaintiff's attorney,		
If you fail to respond, judgment by default will be entered against you for the relief demanded in the complaint. You also must file your answer or motion with the court.					
MAY 1 4 20 Date:	12		CLERK OF COURT Signature of Clerk or Deputy Clerk		

PROOF OF SERVICE

(This section should not be filed with the court unless required by Fed. R. Civ. P. 4 (1))

was re	ceived by me on (date)	•		
	☐ I personally served	the summons on the individual at (p	lace)	
			on (date)	
	☐ I left the summons	at the individual's residence or usua	l place of abode with (name)	
	, a person of suitable age and discretion who resides there,			
	on (date), and mailed a copy to the individual's last known address; or			
	☐ I served the summo	ons on (name of individual)		, who is
	designated by law to accept service of process on behalf of (name of organization)			
			on (date)	_ ; or
	☐ I returned the summ	nons unexecuted because		; or
	Other (specify):			
	My fees are \$	for travel and \$	for services, for a total of \$	0.00
	I declare under penalt	y of perjury that this information is t	rue.	
Date:				*
Duto.			Server's signature	·
			Printed name and title	· · · · · · · · · · · · · · · · · · ·
			Server's address	<u>.</u>