

IN THE SUPERIOR COURT OF THE STATE OF DELAWARE

IN RE ZANTAC (RANITIDINE)
LITIGATION

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C.A. NO. N22C-09-101 ZAN

Submitted: March 7, 2024

Decided: May 31, 2024

OMNIBUS ORDER
ON MOTIONS TO EXCLUDE EXPERT OPINIONS

I. INTRODUCTION

Nearly 75,000 Plaintiffs seek to be heard in Delaware for claims alleging that their cancer was caused due to the ingestion of a heartburn medication commonly known as Zantac. In this early stage of these proceedings, before the Court for disposition are the parties’ competing motions to exclude expert testimony pursuant to Rule 702 of the Delaware Rules of Evidence and *Daubert v. Merrell Dow Pharm. Inc.*,¹ (the “Motions”).

The Motions were the subject of discovery, a three-day “*Daubert*” hearing, multiple layers of briefing, and post-hearing submissions all supported by more than

¹ 509 U.S. 579 (1993).

forty volumes of exhibits. Having considered the pleadings, oral arguments, supplemental submissions, and the full record herein, for the reasons now stated, the parties' Motions are **DENIED**.

II. FACTUAL AND PROCEDURAL HISTORY²

This case involves a molecule known as ranitidine. Ranitidine is marketed under the label name of Zantac. N-Nitrosodimethylamine ("NDMA") is found in ranitidine.³ NDMA causes cancer.⁴

Zantac is a part of a class of medications known as Histamine-2 Receptor Antagonists ("H2Ras").⁵ Ranitidine is a histamine-2 receptor blocker used to "treat heartburn and many other gastro-intestinal disorders, including duodenal ulcers, gastroesophageal reflux disease ("GERD") and esophagitis."⁶

In 1983, based on extensive testing, including humans, the FDA approved ranitidine for prescription use to treat ulcers and later approved it to treat other

² The recitation of the history and facts in this section are for context only.

³ See Plaintiffs' Opposition to Defendants' Motion to Exclude General Causation Experts' Opinions at 14, Trans. ID 71670509 (Dec. 20, 2023) (herein "Pls.' Opp'n to Defs.' Mot. to Exclude Gen. Causation Experts' Op.>").

⁴ *Id.*

⁵ Defendants' Brief in Support of Brand Defendants' and Patheon's Motion to Exclude Plaintiffs' General Causation Experts' Opinions at 3, Trans. ID 71408977 (Nov. 15, 2023) (herein "Defs.' Br. in Supp. of Brand Defs.' & Patheon's Mot. to Exclude Pls.' Gen. Causation Experts' Op.>").

⁶ *In re Zantac*, 644 F. Supp.3d 1075, 1095 (S.D. Fla. 2022).

stomach and esophageal conditions.⁷ In 1995, the FDA authorized ranitidine for over-the-counter (“OTC”) use.⁸ By 2004, the FDA had further approved higher dosages of ranitidine for OTC use.⁹

Zantac was on the market for more than 35 years.¹⁰ During approximately four decades of marketing, there were four brand pharmaceutical companies and generic manufacturers that sold versions of the product.¹¹ GlaxoSmithKline (“GSK”) developed the medication and initially marketed it in prescription form.¹² In 1995, GSK marketed it as an OTC in a joint venture with a predecessor of Pfizer.¹³ In 1998, GSK transferred its rights to sell OTC Zantac in the U.S. to that Pfizer predecessor.¹⁴ In 2006, Defendant Boehringer Ingelheim (“BI”) acquired the rights to sell OTC Zantac.¹⁵ In 2017, Defendant Sanofi began selling OTC Zantac after acquiring the brand from BI.¹⁶

⁷ Defs.’ Br. in Supp. of Brand Defs.’ & Patheon’s Mot. to Exclude Pls.’ Gen. Causation Experts’ Op. at 4.

⁸ *Id.*

⁹ *Id.*

¹⁰ *Id.*

¹¹ *Id.*

¹² Defs.’ Br. in Supp. of Brand Defs.’ & Patheon’s Mot. to Exclude Pls.’ Gen. Causation Experts’ Op. at 4, n1.

¹³ *Id.*

¹⁴ *Id.*

¹⁵ *Id.*

¹⁶ *Id.*

In September of 2019, Valisure, an online pharmacy submitted a citizen petition to the FDA claiming detection of “extremely high levels of N-Nitrosodimethylamine (NDMA)” in ranitidine.¹⁷ Valisure reported NDMA at levels in excess of three million nanograms per tablet. This far exceeded the limit of 96 nanograms per day that the FDA had set for NDMA ingestion in the context of an unrelated class of medications.¹⁸

After reviewing Valisure’s petition, the FDA raised concerns about the testing methodology.¹⁹ FDA and ranitidine manufacturers studied NDMA in ranitidine and examined whether ranitidine use increases cancer risks in patients.²⁰ Over the next month, some tests revealed amounts lower than what Valisure reported, and some lots tested revealed amounts below the acceptable daily intake (ADI).²¹

In September and October of 2019, then-existing ranitidine manufactures recalled their products. And by April 2020—after further testing confirmed NDMA levels in some samples continued to exceed ADI—the FDA requested manufactures initiate a market withdrawal of all remaining batches then remaining on the market.²² After the recall of ranitidine-containing Zantac, litigation ensued around the country.

¹⁷ Defs.’ Br. in Supp. of Brand Defs.’ & Patheon’s Mot. to Exclude Pls.’ Gen. Causation Experts’ Op. at 5.

¹⁸ *Id.*

¹⁹ *Id.*

²⁰ *Id.*

²¹ *Id.* (citing FDA, *FDA Statement: Statement on new Testing Results, Including Low Levels of Impurities in Ranitidine Drugs* (2019) (Brown Decl. Ex. 89) at 1).

²² *Id.* at 8.

A. NATIONAL PROCEDURAL HISTORY - THE “MDL”²³

To address these claims, on February 6, 2020, the United States Judicial Panel on Multidistrict Litigation established a multidistrict litigation process (the “MDL”) in the U.S. District Court for the Southern District of Florida in West Palm Beach for all pretrial purposes. The Panel ordered federal lawsuits for personal injury and economic damages from the purchase or use of Zantac to be transferred to the MDL.

As part of MDL management, a Census Registry (“The Registry”) was created to allow the parties and the Court to “understand the nature of the unfiled claims that are a part” of the MDL.²⁴ The MDL Court held a *Daubert* hearing in March 2018.

On December 6, 2022, the MDL Court issued its opinion on *Daubert* and summary judgment motions (“MDL Order”).²⁵ In its 200-page opinion, the MDL Court, in pertinent part, excluded those plaintiffs’ experts’ general causation opinions and granted summary judgment for Defendants.²⁶

B. LITIGATION IN OTHER STATES

Similar suits were also proceeding in state courts throughout the United States. The largest one, other than here, was a coordinated proceeding in California, the

²³ See generally, *In re Zantac*, 644 F.Supp.3d at 1095.

²⁴ *Id.* at 1096.

²⁵ As of the date of this ruling, it appears that the MDL decision is on appeal in the Eleventh Circuit Court of Appeals.

²⁶ See generally, *id.*

Judicial Council Coordinated Proceeding (“JCCP”). In the California state court, several thousand cases were being coordinated in the JCCP, with sixteen bellwether trials scheduled for 2024. The JCCP Plaintiffs were pursuing their claims for the same cancers claimed here. Those cases advanced beyond the general causation phase.²⁷

C. THE PARTIES AND PROCEDURAL HISTORY HERE

Plaintiffs in this litigation were not before the federal MDL Court. Nor are the experts the same. Plaintiffs here are pursuing ten cancers—bladder, esophageal, gastric, liver, pancreatic, breast, colorectal, kidney, lung and prostate. Notably, in the MDL, Plaintiffs’ Leadership (also not present here) notified that Court that it had decided not to pursue general causation expert reports for breast and kidney cancers and initially narrowed their list from ten to eight cancers.²⁸ In January of 2022, they again notified the MDL Court that they were not moving forward with certain cancers, and again, narrowed the list to five cancers. Thus, five of the cancer claims here were not before the MDL Court.²⁹

In September of 2022, nearly 75,000 complaints were filed in this Court.

²⁷ See Pls.’ Opp’n to Defs.’ Mot. to Exclude Gen. Causation Experts’ Op. Ex 75.

²⁸ *In re Zantac*, 644 F.Supp.3d at 1098.

²⁹ *Id.* (The MDL noted in its final disclosure that the plaintiffs in the MDL intended to “prove that ranitidine causes bladder, esophageal, gastric, liver, and pancreatic cancers (the ‘Designated Cancers’), as opposed to other cancers (‘Non-Designated Cancers.’). The Defendants do not address Non-Designated Cancers in the *Daubert* motions, so individual cases in which Plaintiffs allege their ranitidine use caused their Non-Designated Cancers remain pending at this time and are not the subject of this Order.”).

Plaintiffs allege Defendants collectively bear responsibility for their cancer diagnoses, and the related injuries or deaths caused from their ingestion of the medication known as Zantac.

The Defendants are GlaxoSmithKline LLC (“GSK”), Boehringer Ingelheim Pharmaceuticals, Inc., Boehringer Ingelheim Corporation, and Ingelheim U.S.A. Corporation (collectively, B.I.), Sanofi US Services Inc., Sanofi-Aventis U.S. LLC, and Chattem, Inc. (collectively, “Sanofi), Pfizer Inc. (“Pfizer”) (together with all those just mentioned are referred to as the “Brand Defendants”) and Patheon, (all collectively “Defendants”).³⁰

In these initial proceedings, the first phase addresses “general causation,” which involves the question of whether the ingestion of this product is capable of causing cancer as alleged. To carry their burden at this stage, Plaintiffs have retained ten experts to offer opinions on general causation for the ten mentioned cancers. Defendants move to exclude them all.³¹ Plaintiffs also move to exclude certain opinions proffered by Defendants’ sole General Causation Expert, William C.

³⁰ Brand Defendants’ and Patheon’s Motion to Exclude Plaintiffs’ Expert Dr. Charles Jameson, Trans. ID 71409144 (Nov. 15, 2023) (herein “Brand Defs.’ & Patheon’s Mot. to Exclude Pls.’ Expert Dr. Jameson”).

³¹ Defendants move to exclude all of Plaintiffs’ General Causation Experts: Drs. Charles William Jameson, PhD; William Sawyer, PhD; Alfred I. Neugut, M.D., PhD; Vinod K. Rustgi M.D., MBA; Ioannis Hatzaras, M.D. MPH, PhD, F.A.C.S.; Dan J. Raz, M.D.; Bruce J. Trock, MPH, PhD; George Miller, M.D.; Pablo Leone, M.D.; and Vitaly Margulis, M.D. (collectively “Plaintiffs’ General Causation Experts”).

Zamboni, Pharm.D., PhD.

III. APPLICABLE LEGAL STANDARDS

As in any products liability case, each party bears the burden of proof on the admissibility of their expert opinion testimony.³² Delaware Rule of Evidence 702 addresses the admissibility of expert testimony. It provides:

A witness who is qualified as an expert by knowledge, skill, experience, training, or education may testify in the form of an opinion or otherwise if:

- (a) the expert’s scientific, technical, or other specialized knowledge will help the trier of fact to understand the evidence or to determine a fact in issue;
- (b) the testimony is based on sufficient facts or data;
- (c) the testimony is the product of reliable principles and methods; and
- (d) the expert has reliably applied the principles and methods to the facts of the case.³³

This rule is nearly identical to Federal Rule of Evidence 702.³⁴ Our Supreme Court has interpreted Federal Rule of Evidence 702, addressed the admissibility of expert testimony,³⁵ and adopted the holdings of *Daubert v. Merrell Dow Pharm.*,

³² *Minner v. Amer. Mort. & Guar. Co.*, 791 A.2d 826, 843 (Del. Super. 2000).

³³ D.R.E. 702.

³⁴ *Barrera v. Monsanto Co.*, 2019 WL 2331090, at *3 (Del. Super. May 31, 2019).; *see also* *Minner*, 791 A.2d at 833 n.2.

³⁵ *See Tumlinson v. Advanced Micro Devices, Inc.*, 81 A.3d 1264, 1269 (Del. 2013) (herein “*Tumlinson*”).

Inc. (“*Daubert*”) and its progeny as the correct interpretation of Delaware Rule of Evidence 702.³⁶

A. THE *DAUBERT* STANDARD

Decided in 1993 by the United States Supreme Court, *Daubert* is a name well-known to most lawyers. Even so, the Court has urged litigants to appreciate “the perspective from which this Court will view so-called *Daubert* motions, and the process by which such motions will be addressed.”³⁷ This encouragement merits respect, as *Daubert* at first blush may seem contradictory, expanding the power of the trial court by rejecting the “general acceptance” requirement for admissibility while at the same time emphasizing the limitations on the court’s role in deference to the role of the jury.³⁸ And although *Daubert* rejected the exclusivity of the “general acceptance” requirement, an expert’s “access to the courtroom is not unfettered.”³⁹

A few factors can be implicated in *Daubert* reviews. Some are enumerated in Rule 702; others, some identified *in Daubert*, are deemed nonexclusive.⁴⁰ But even the expanded list is not exclusive. None of the *Daubert* factors, specific or

³⁶ See *M.G. Bancorporation, Inc. v. LeBeau*, 737 A.2d 513, 522 (Del. 1999).

³⁷ *In re Asbestos Litig.*, 911 A.2d 1176, 1197 (Del. Super. 2006).

³⁸ *Minner*, 791 A.2d at 841.

³⁹ *In re Asbestos Litig.*, 911 A.2d at 1197.

⁴⁰ *Daubert*, 509 U.S. at 590.

otherwise, is binding on the trial court.⁴¹ The trial court also has broad discretion to consider factors not articulated by *Daubert*.⁴²

B. THE GATEKEEPER AND THE JURY

Predominant among *Daubert*'s holdings is the recognition of the trial court's role as gatekeeper. In that role, the "Trial Judge...insure[s] that the scientific testimony is not only relevant but reliable."⁴³ While *Daubert* may require trial courts to dive deeper into certain preliminary facts than had historically been the case, it was not intended to abrogate the jury's constitutionally protected role as the ultimate fact-finder; a role the courts of this State defend vigorously.⁴⁴ *Daubert* and its progeny have repeatedly emphasized the importance of the trial system and the role of juries as the ultimate arbiters in expert evidentiary issues.

Therefore, "[a]s a threshold matter, *Daubert* neither requires nor empowers Trial Courts to determine which of . . . competing scientific theories has the best performance."⁴⁵ *Daubert* "requires only that the trial court determine whether the

⁴¹ *Tumlinson*, 81 A.3d at 1269.

⁴² *Id.* at 1272–73; *Long v. Weider Nutrition Group*, 2004 WL 1543226, at *5 (Del. Super. June 25, 2004) (citing *Kumho Tire Co., Ltd. v. Carmichael*, 526 U.S. 137, 150 (1999)).

⁴³ *Minner*, 791 A.2d at 843 (citations omitted).

⁴⁴ *In re Asbestos Litig.*, 911 A.2d at 1199; *see. e.g., Kaur v. Bos. Sci. Corp.*, 2022 WL 1486178, at *3 (Del. Super. May 11, 2022) (an expert's investigations and assumptions "are readily subject to cross examination and to evaluation by the fact finder . . .").

⁴⁵ *Minner*, 791 A.2d at 848 (citation omitted).

proponent of the evidence has demonstrated that scientific conclusions have been generated using sound and reliable approaches.”⁴⁶ But “the judge is not a scientist and the courtroom is not a science laboratory.”⁴⁷ Indeed,

“[i]t would be unreasonable to conclude that the subject of scientific testimony must be ‘known’ to a certainty [N]ot knowing the mechanism whereby a particular agent causes a particular effect is not always fatal to a plaintiff’s claim. Causation can be proved even where we do not know precisely *how* the damage occurred if there is sufficiently compelling proof that the agent must have caused the damage *somehow*.”⁴⁸

As such, to determine the admissibility of scientific evidence consistent with *Daubert*, the trial judge must determine whether:

- (1) the witness is qualified as an expert by knowledge, skill, experience, training or education;
- (2) the evidence is relevant;
- (3) the expert’s opinion is based upon information reasonably relied upon by the experts in the particular field;
- (4) the expert testimony will assist the trier of fact to understand the evidence or to determine a fact in

⁴⁶ *In re Asbestos*, 911 A.2d at 1201; *see also Minner*, 791 A.2d at 842 (citations omitted).

⁴⁷ *Daubert*, 509 U.S. at 596–97 (quoting “Introduction to Reference Manual on Scientific Evidence”, Fed. Jud. Ctr. at 2 (2000)).

⁴⁸ *Daubert v. Merrell Dow Pharms., Inc.*, 43 F.3d 1311, 1314 (9th Cir. 1995) (emphasis in original); *accord, Bowen v. E.I. Du Pont de Nemours and Co.*, 2005 WL 1952859, at *9 (Del. Super. June 23, 2005) (herein “*Bowen*”), *aff’d* 906 A.2d 787 (Del. 2006) (explaining that the trial court need not “decide the admissibility of scientific evidence with the degree of certainty required in scientific circles”).

issue; and

- (5) the expert will not create unfair prejudice or confuse or mislead the jury.⁴⁹

The gatekeeper must apply these particular “factors in a flexible manner that takes into account the particular specialty of the expert under review and the particular facts of the underlying case.”⁵⁰ “Where the question of admissibility is a close one, exclusion of the evidence is not appropriate where cross examination, the presentation of contrary evidence and careful instruction regarding the burden of proof will insure that the jury is not misled or confused.”⁵¹ Restated, “[t]he reliability requirement is not a tool for the Court to use to exclude questionably reliable evidence.”⁵² This Court’s refusal to establish a bright line rules for proving causality has previously been considered.⁵³ And no doubt, “the requisite proof necessary to establish causation will vary greatly case by case.”⁵⁴

The Supreme Court in *Daubert* was more direct: “Vigorous cross-examination, presentation of contrary evidence, and careful instruction on the burden of proof are the traditional and appropriate means of attacking shaky but admissible

⁴⁹ *Bowen v. E.I. DuPont de Nemours & Co., Inc.*, 906 A2d 787, 795 (Del. 2006) (herein “*Bowen I*”).

⁵⁰ *Id.*; *Scaife v. Astrazeneca LP*, 2009 WL 1610575, at *14 (Del. Super. June 9, 2009).

⁵¹ *Bowen*, 2005 WL 1952859, at *8, *aff’d sub nom. Bowen II*, 906 A.2d at 787.

⁵² *Barrera*, 2019 WL 2331090, at *10.

⁵³ *Id.* at *5 (citing *In re Zoloft*, 858 F.3d 787, 783 (3d Cir. 2017)).

⁵⁴ *In re Zoloft*, 858 F.3d at 787.

evidence.”⁵⁵ Thus guided, courts confronted by “shaky but admissible evidence” conduct their *Daubert* analyses “with a ‘liberal thrust’ favoring admission.”⁵⁶

C. DAUBERT AND GENERAL CAUSATION CONSIDERATIONS

As the preliminary question concerns *general causation*, this Court finds the federal district court’s approach in *In re Roundup Products Liability Litigation*⁵⁷ instructive:

The question at this early phase in the proceedings - the ‘general causation phase - is whether a reasonable jury could conclude that glyphosate . . . can cause Non Hodgkins Lymphoma (“NHL”) There are two significant problems with the plaintiffs’ presentation, which combine to make this a very close question.

* * *

The evidence, viewed in its totality, seems too equivocal to support any firm conclusion that glyphosate causes NHL. This calls into question the credibility of some of the plaintiffs’ experts, who have confidently identified a causal link.

However, the question at this phase is not whether the plaintiffs’ experts are right. The question is whether they have offered opinions that would be admissible at a jury trial. And the case law - particularly Ninth Circuit case law - emphasizes that a trial judge should not exclude an expert opinion merely because he thinks it’s shaky, or

⁵⁵ *Daubert*, 509 U.S. at 596.

⁵⁶ *Messick v. Novartis Pharmaceuticals Corp.*, 747 F.3d 1193, 1196 (9th Cir. 2014) (quoting *Daubert*, 509 U.S. at 588); *M.G. Bancorporation, Inc.*, 737 A.2d at 522.

⁵⁷ 390 F.Supp.3d 1102 (N.D. CA 2018) (herein “*In re Roundup*”).

results.”⁴⁹⁰ “With this information, Defendants are able to do their own analysis, their own processing and integration, report their results, and do their own calculations of means, minimum and maximum values, or any other statistical measure they wish. They did no such thing.”⁴⁹¹ Defendants resort to *ipse dixit*: the production, characterized as they deem appropriate, makes reproduction impossible. They never say they tried to reproduce Emery’s results. Instead, they cite to the MDL’s Order.⁴⁹²

Defendants devote considerable space to the numerous failings and inconsistencies they find in Emery’s MDL and Delaware Reports.⁴⁹³ They also chastise Emery for its inability to reproduce its own results. This criticism might carry more weight if Emery were somehow prohibited from updating the report, or even more so if Emery had not explained to Defendants why the data was updated in the first instance.⁴⁹⁴ Further, none of the changes cited by Defendants concerns the baseline testing results.⁴⁹⁵ Also, some of the results of the changes made by Dr. Cheu following his review and re-processing are “actually lower than those reported

⁴⁹⁰ *Id.* at 269.

⁴⁹¹ *Id.* at 273.

⁴⁹² Defs.’ Br. in Supp. of Brand Defs.’ & Patheon’s Mot. to Exclude Emery Pharma at 56–58 (citing *In re Zantac*, 644 F.Supp.3d at 1130).

⁴⁹³ *See id.* at 57–63.

⁴⁹⁴ *See* Pls.’ Opp’n to Defs.’ Mot. to Exclude Gen. Causation Experts’ Op. at 271–74.

⁴⁹⁵ *Id.* at 276.

In Delaware, our jurisprudence counsels that, subject to earnest deliberation, trial courts entrust questions of science to the scientists. Here, opposing teams of highly educated, skilled expert medical witnesses offer competing opinions. Through well-trained counsel, their efforts only clarify the distinct opposition that defines their respective positions. It would be improper to simply dismiss these experts as “poseurs or witnesses for hire. They are serious scientists.”⁵¹⁶ As gatekeeper, the Court has found that each side has carried its required burden of demonstrating the reliability of its proffered Rule 702 evidence. Any remaining challenges will be made at trial via cross-examination and introduction of counter evidence. Having considered the full record herein, the parties’ *Daubert* challenges fail and their Motions to Exclude are **DENIED**.

IT IS SO ORDERED.

/s/ Vivian L. Medinilla
Vivian L. Medinilla
Judge

cc: All Counsel of Record

⁵¹⁶ *In re Asbestos Litig.*, 911 A.2d at 1207.