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13 *Attorneys for Plaintiffs Tirrell Allen and*  
14 *LaToya Allen*

*Attorneys for Defendants Global Blood*  
*Therapeutics, Inc. and Pfizer Inc.*

15  
16 **UNITED STATES DISTRICT COURT**  
17 **NORTHERN DISTRICT OF CALIFORNIA**  
18 **SAN FRANCISCO DIVISION**

19 TIRRELL ALLEN and LATOYA  
20 ALLEN,

21 Plaintiffs,

22 v.

23 GLOBAL BLOOD THERAPEUTICS,  
24 INC. and PFIZER INC.,

25 Defendants.  
26  
27  
28

Case No. 3:24-cv-07786-TLT

**JOINT CASE MANAGEMENT  
STATEMENT**

Date: August 21, 2025  
Time: 2:00 p.m.  
Place: Remote (Zoom)  
Judge: Hon. Trina L. Thompson

**JOINT CASE MANAGEMENT STATEMENT**

Pursuant to Civil Local Rule 16-9, the Standing Order for All Judges of the Northern District of California regarding Contents of Joint Case Management Statement, and this Court’s Standing Order for Civil Cases, Plaintiffs Tirrell Allen and LaToya Allen (“Plaintiffs”) and Defendants Global Blood Therapeutics, Inc. and Pfizer Inc. (“Defendants”) (collectively, “the Parties”), hereby submit the following joint statement.

**1. Jurisdiction and Service**

Plaintiff Tirrell Allen filed his original Complaint on November 7, 2024 (Dkt. 1) and served Defendants on January 6, 2025 (Dkts. 18, 19). This Court has subject matter jurisdiction over this action pursuant to 28 U.S.C. §§ 1331 and 1367. No issues exist regarding personal jurisdiction or venue, and no Defendant remains unserved.

**2. Facts****a. Plaintiffs’ Statement**

This is an action for damages related to Defendants’ conduct in connection with the development, design, testing, manufacturing, labeling, packaging, promoting, advertising, marketing, distribution, and selling of Oxbryta (generic name: voxelotor), a prescription medication used to treat sickle cell disease. The FDA approved Oxbryta under the accelerated approval pathway in 2019 for the treatment of sickle cell disease in adults and pediatric patients 12 years of age and older. In 2021, FDA granted accelerated approval of Oxbryta for the treatment of sickle cell disease in patients 4 to 11 years of age. Accelerated approval is based on a surrogate or intermediate clinical endpoint that is reasonably likely to predict clinical benefit, allowing for earlier approval of drugs that treat serious conditions and fill an unmet medical need. In general, FDA requires post-marketing studies to verify and describe the clinical benefit of medications approved under this program. Defendants marketed Oxbryta through various forms of media and promised its purchasers would “experience less sickling.”

On September 25, 2024, Defendants announced they were voluntarily withdrawing the medication from the market, ceasing distribution, and discontinuing all active clinical trials and expanded access programs for Oxbryta “because recent data indicate the benefit of Oxbryta does not

1 outweigh the risks for the sickle cell patient population.” Defendants noted that their decision was  
2 “based on the totality of clinical data that now indicates the overall benefit of Oxbryta no longer  
3 outweighs the risk in the approved sickle cell patient population. The data suggest an imbalance in  
4 vaso-occlusive crises and fatal events which require further assessment.”

5 Plaintiff Tirrell Allen is a 43-year old male who was diagnosed with sickle cell disease as a  
6 child. While on Oxbryta, he experienced an increased rate of vaso-occlusive crises (VOCs), suffered  
7 a stroke, and was hospitalized.

#### 8 **b. Defendants’ Statement**

9 This case is about Oxbryta (voxelotor), a prescription medicine developed by Global Blood  
10 Therapeutics, Inc. (“GBT”) for the treatment of sickle cell disease (“SCD”). SCD is a lifelong,  
11 inherited disease that affects hemoglobin, the protein in red blood cells that is responsible for  
12 delivering oxygen throughout the body. It affects approximately 100,000 people in the United  
13 States. In patients with sickle cell disease, abnormal hemoglobin causes red blood cells to become  
14 rigid, sticky, and “sickle”-shaped. These sickled red blood cells clump together and restrict the flow  
15 of oxygen, causing pain events called vaso-occlusive crises (“VOCs”), acute chest syndrome,  
16 swelling, anemia, and strokes, among other complications.

17 In 2019, the FDA approved Oxbryta for use by adults and pediatric patients 12 years and  
18 older, based on clinical trial results as well as the significant unmet medical needs of patients with  
19 sickle cell disease; two years later, the agency expanded the medication’s approved use to patients  
20 as young as 4 years old. Oxbryta was the first approved sickle cell treatment to target the root cause  
21 of sickle cell disease; by improving the ability of hemoglobin to bind to oxygen, the medicine helps  
22 red blood cells maintain their normal shape. In a clinical trial, patients treated with Oxbryta  
23 demonstrated a statistically significant improvement in hemoglobin response, and showed no  
24 increase in vaso-occlusive crises.<sup>1</sup>

25 Pfizer Inc. (“Pfizer”) acquired GBT in October 2022, and continued to study the benefit of  
26 Oxbryta in both confirmatory studies and real-world registries. In September 2024, Pfizer

27 \_\_\_\_\_  
28 <sup>1</sup> Center for Drug Evaluation & Research, App No. 213137, Multi-Discipline Review & Evaluation (Division Director  
Summary Review for Regulatory Action at 12), *available at*  
[https://www.accessdata.fda.gov/drugsatfda\\_docs/nda/2019/213137Orig1s000Multidiscipline.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/nda/2019/213137Orig1s000Multidiscipline.pdf).

1 announced the voluntary withdrawal of Oxbryta following an initial review of available data from  
2 post-marketing and registry-based studies, which appeared to show an unexpectedly higher rate of  
3 VOCs in some Oxbryta patients, and a higher number of deaths among some patients taking Oxbryta  
4 for a longer period of time. Pfizer notified the FDA and other regulatory authorities that it was  
5 continuing to review all available data regarding Oxbryta; that analysis is ongoing.

6 Approximately six weeks later, on November 7, 2024, Plaintiff filed his Complaint in this  
7 action, alleging that, during the one-month period he was taking Oxbryta, it caused him to experience  
8 a “higher rate of VOCs” than prior to taking the medication, and a stroke for which he was  
9 hospitalized.

### 10 **3. Legal Issues**

11 Plaintiffs’ Amended Complaint alleges nine claims: (1) Strict Products Liability – Design  
12 Defect; (2) Strict Products Liability – Failure to Warn; (3) Negligence; (4) Negligent  
13 Misrepresentation; (5) Breach of Express Warranties; (6) Breach of Implied Warranties; (7 )Quasi-  
14 Contract; (8) Violation of the Illinois Consumer Fraud and Deceptive Business Practices Act, 815  
15 Ill. Comp. Stat. 505/1 - 505/12; and (9) loss of consortium.

#### 16 **a. Plaintiff’s Statement**

17 Plaintiff maintains that Defendants are liable based on the causes of action listed above and  
18 preliminarily identify the following legal issues: whether Defendants shall be held liable under  
19 Plaintiff’s theories of recovery; Whether Defendant’s conduct rises to the level of punitive damages;  
20 whether Defendant’s advertisements violate California law by being false and/or deceptive;

#### 21 **b. Defendants’ Statement**

22 Defendants dispute Plaintiff’s allegations, deny that they are liable for any of the claims  
23 asserted by Plaintiff in the Complaint, and, at the appropriate time, will file an answer with  
24 affirmative defenses. The principal legal issues include, but are not limited to: whether the  
25 Complaint should be dismissed for Plaintiff’s failure to state a claim; whether Plaintiff’s strict  
26 liability claims are recognized under California law; whether any alleged defect in Oxbryta caused  
27 or contributed to Plaintiff’s claimed injuries; whether the warnings for Oxbryta were adequate;  
28 whether Plaintiff’s claims are barred by the learned intermediary doctrine; whether Plaintiff’s claims

1 are barred by federal preemption; whether Plaintiff has standing to pursue injunctive relief for his  
2 claims; whether Defendants' alleged failure to warn caused Plaintiff's injuries; and whether Plaintiff  
3 relied on any statements or warranties about Oxbryta.

4 **4. Motions**

5 Defendants filed a motion to dismiss the amended complaint on April 23, 2025 (Dkt. 47).  
6 That motion is fully briefed, and set for hearing on October 14, 2025.

7 There are no other pending motions. The Parties reserve the right to file other motions as  
8 appropriate, including motions for summary judgment (or partial summary judgment), and pretrial  
9 motions, including motions *in limine*.

10 **5. Amendment of Pleadings**

11 Plaintiffs filed an amended complaint on March 12, 2025. Pursuant to the Court's Case  
12 Management and Scheduling Order (Dkt. 32), the last day to amend pleadings was May 22, 2025.

13 **6. Evidence Preservation**

14 The Parties certify that they have reviewed the Guidelines Relating to the Discovery of  
15 Electronically Stored Information, and confirm that they have met and conferred pursuant to Fed. R.  
16 Civ. P. 26(f) regarding reasonable and proportionate steps taken to preserve evidence relevant to the  
17 issues reasonably evident in this action. The Parties are aware of and complying with their  
18 preservation obligations, and will advise the Court in the event they are unable to reach an agreement  
19 on ESI-related issues.

20 **7. Disclosures**

21 Neither party has exchanged initial disclosures as of the date of the filing of this Joint Case  
22 Management Statement. The Parties propose that they exchange their Initial Disclosures within 30  
23 days after the Court rules on Defendants' motion to dismiss the amended complaint.

24 **8. Discovery**

25 **a. Discovery Taken to Date**

26 Plaintiffs served initial discovery requests on Defendants on May 15, 2025.  
27  
28

1                   **b. Scope of Anticipated Discovery**

2                           **i. Plaintiff's Statement**

3                   Plaintiffs intend to seek discovery from Defendants and third party sources related to the  
4 following topics, among other things, a) all study data that led to the Oxbryta recall, b) Defendant  
5 Pfizer's acquisition and current relationship with Defendant Global Blood Therapeutics, c) adverse  
6 event reporting data, d) European Medicine Agency Study GBT440-032 and Study GBT440-042  
7 data, e) summary basis of approval for application for Oxbryta and f) information related to  
8 Defendant's development, design, testing, manufacturing, labeling, packaging, promoting,  
9 advertising, marketing, distribution, and selling of Oxbryta.

10                           **ii. Defendants' Statement**

11                   If this case proceeds to discovery, Defendants intend to seek discovery from Plaintiff and  
12 third parties regarding, among other topics: (a) Plaintiff's past and ongoing medical evaluation and  
13 treatment; (b) the decision of Plaintiff's healthcare providers to prescribe Oxbryta to Plaintiff; (c)  
14 details concerning Plaintiff's ingestion of Oxbryta; (d) how and when Plaintiff learned of the alleged  
15 relationship between his ingestion of Oxbryta and his alleged injuries; (e) Plaintiff's alleged injuries  
16 and his support for his assertions that Oxbryta caused those injuries; (f) warnings, labels, and other  
17 promotional materials about Oxbryta, if any, that Plaintiff relied upon; and (g) Plaintiff's damages.

18                           **c. Modifications to the Discovery Rules**

19                   The Parties do not currently request any modifications to the Discovery Rules but reserve  
20 the right to request modifications as the litigation proceeds.

21                           **d. Agreement to Enter a Stipulated E-Discovery Order**

22                   The Parties agree to cooperate and work in good faith toward reaching an agreement on a  
23 stipulation regarding the preservation and production of electronically stored information, as well as  
24 a protective order governing the discovery and use of confidential information. If agreement cannot  
25 be reached, the Parties will seek the Court's assistance.

26                           **e. Discovery Disputes**

27                   The Parties have not identified any discovery disputes at this time.

28                   **9. Class Actions**

1 The Plaintiff does not assert claims on behalf of a class.

2 **10. Related Cases**

3 **a. Federal Court**

- 4 1. *Jolly, et al. v. Global Blood Therapeutics, Inc. & Pfizer Inc.*, Case No.  
5 3:24-CV-09345-TLT (N.D. Cal.). Defendants filed a motion to  
6 dismiss the first amended complaint on April 23, 2025. That motion  
7 is fully briefed and set for hearing on October 14, 2025. A mediation  
8 is scheduled on September 9, 2025.
- 9 2. *Frazier v. Global Blood Therapeutics, Inc. & Pfizer Inc.*, Case No.  
10 3:25-cv-04027-TLT (N.D. Cal.). The complaint was filed on May 8,  
11 2025, and served on Defendants on July 1, 2025. Defendants filed a  
12 motion to dismiss on July 22, 2025, which is set for hearing on  
13 October 14, 2025.
- 14 3. *Ford v. Global Blood Therapeutics, Inc. & Pfizer Inc.*, Case No. 3:25-  
15 cv-04229-TLT (N.D. Cal.). The complaint was filed on May 16, 2025,  
16 and served on Defendants on July 1, 2025. Defendants filed a motion  
17 to dismiss on July 22, 2025, which is set for hearing on October 14,  
18 2025.
- 19 4. *Marvin Cosey, as the Guardian of Amiyah S. Cosey v. Global Blood*  
20 *Therapeutics, Inc. & Pfizer Inc.*, Case No. 1:25-CV-09400 (N.D. Ill.).  
21 Status: Plaintiff filed the complaint in the Circuit Court of Cook  
22 County, Illinois on June 27, 2025. Plaintiff served the complaint on  
23 Defendant Pfizer Inc. on July 11, 2025, and on Defendant Global  
24 Blood Therapeutics, Inc. on July 14, 2025. On August 8, 2025,  
25 Defendants removed the case to the United States District Court for  
26 the Northern District of Illinois. On August 12, 2025, Defendants filed  
27 an uncontested motion for extension of time to file a responsive  
28

1 pleading to September 12, 2025, which was granted on August 14,  
2 2025.

3 **b. State Court**

4 There are currently fourteen product liability cases pending in state courts (including nine in  
5 California) that have been served on Pfizer and/or GBT involving claims that plaintiffs suffered  
6 personal injuries from consuming Oxbryta. Those cases are:

- 7 1. *Tolulope Afolabi v. Pfizer Inc., Global Blood Therapeutics, Inc., & Does 1*  
8 *through 100*, Case No. 24-CIV-08331 (Cal. Super. Ct. – San Mateo Cnty.)  
9 Status: Defendants filed a demurrer to Plaintiff’s complaint on March 17,  
10 2025, and further briefing on the demurrer is in progress. A hearing on  
11 Defendants’ demurrer is set for January 15, 2026. The next case management  
12 conference is on September 3, 2025. Plaintiff has served initial discovery  
13 requests.  
14
- 15 2. *Raven Favor v. Global Blood Therapeutics, Inc.*, Case No. 25-CIV-01314  
16 (Cal. Super. Ct. – San Mateo Cnty.) Status: Defendant filed a demurrer to  
17 Plaintiff’s complaint on April 21, 2025, and further briefing on the demurrer  
18 is in progress. A hearing on Defendant’s demurrer is set for December 4,  
19 2025. The next case management conference is on September 3, 2025.  
20
- 21 3. *Asja Joseph v. Global Blood Therapeutics, Inc.*, Case No. 25-CIV-01315  
22 (Cal. Super. Ct. – San Mateo Cnty.) Status: Defendant filed a demurrer to  
23 Plaintiff’s complaint on April 21, 2025, and further briefing on the demurrer  
24 is in progress. A hearing on Defendant’s demurrer is set for December 4,  
25 2025. The next case management conference is on September 3, 2025.  
26
- 27 4. *Deborah Majeeda Snead v. Pfizer Inc. & Global Blood Therapeutics, Inc.*,  
28 Case No. 25-CIV-02200 (Cal. Super. Ct. – San Mateo Cnty.) Status:

1 Defendants filed a demurrer to Plaintiff's complaint on May 19, 2025, and  
2 further briefing on the demurrer is in progress. A hearing on Defendants'  
3 demurrer is set for April 23, 2026. The next case management conference is  
4 on September 3, 2025.

5 5. *Trebor Hardiman v. Global Blood Therapeutics, Inc.*, Case No. 25-CIV-  
6 03836 (Cal. Super. Ct. – San Mateo Cnty.) Status: Plaintiff served the  
7 complaint—originally filed in the Superior Court of San Francisco County—  
8 on Defendant Global Blood Therapeutics, Inc. on November 4, 2024. A joint  
9 stipulation for change of venue to the Superior Court of San Mateo County  
10 was filed on February 24, 2025. The case was transferred to the Superior  
11 Court of San Mateo County as of May 20, 2025. Plaintiff filed the amended  
12 complaint on July 11, 2025. Defendant filed a demurrer to Plaintiff's amended  
13 complaint on August 8, 2025. The next case management conference is on  
14 September 3, 2025.

15 6. *Marcia Smith v. Global Blood Therapeutics, Inc.*, Case No. 25-CIV-05179  
16 (Cal. Super. Ct. – San Mateo Cnty.). Status: Plaintiff served the complaint—  
17 originally filed in the Superior Court of San Francisco County—on Defendant  
18 Global Blood Therapeutics, Inc. on February 14, 2024. A joint stipulation for  
19 change of venue to the Superior Court of San Mateo County was filed on  
20 March 17, 2025, and received by the court on June 26, 2025. Defendant filed  
21 a demurrer to Plaintiff's complaint on August 11, 2025. An initial case  
22 management conference is set for December 8, 2025.

23 7. *Laura Christine Matteliano-Madu v. Children's Hospital & Research Center*  
24 *at Oakland, et al.*, Case No. 25CV117566 (Cal. Super. Ct. – Alameda Cnty.).  
25 Status: Defendants filed a demurrer to Plaintiff's complaint on July 10, 2025,  
26 which is set for hearing on January 6, 2026. An initial case management  
27 conference is set for September 4, 2025.  
28

- 1 8. *Q.G., et al. v. Pfizer Inc., Global Blood Therapeutics, Inc., & Does 1 through*  
2 *100*, Case No. 25-CIV-04717 (Cal. Super. Ct. – San Mateo Cnty.). Status:  
3 Plaintiffs filed the complaint on June 23, 2025, and served it on Defendant  
4 Global Blood Therapeutics, Inc. on June 25, 2025. Plaintiffs filed the  
5 amended complaint on July 21, 2025. Defendants intend to file a demurrer to  
6 the amended complaint. An initial case management conference is set for  
7 October 10, 2025.
- 8 9. *Reginald Leach v. Global Blood Therapeutics, Inc.*, Case No. 25-CIV-05784  
9 (Cal. Super. Ct. – San Mateo Cnty.). Status: Plaintiff filed the complaint on  
10 July 30, 2025. Plaintiff served the complaint on Defendant Global Blood  
11 Therapeutics, Inc. on August 11, 2025.
- 12 10. *Clarissa M. Cruz, et al. v. Cooper University Hospital, et al.*, CAM-L-  
13 001954-25 (N.J. Super Ct. – Camden Cnty). Status: Plaintiffs filed the  
14 complaint on June 12, 2025. Plaintiffs served the complaint on Defendant  
15 Pfizer Inc. on June 26, 2025.
- 16 11. *Casseus v. Pfizer Inc. & Global Blood Therapeutics Inc.*, Case No.  
17 158036/2025 (N.Y. Sup. Ct. – N.Y. Cnty.). Status: Plaintiff filed the  
18 complaint in the Supreme Court of the State of New York, New York County  
19 on June 24, 2025. Defendants removed the case to the U.S. District Court for  
20 the Southern District of New York on June 27, 2025. Pursuant to a joint  
21 stipulation, the court remanded the case to state court on July 23, 2025.
- 22 12. *Kael Melan Butler v. Pfizer, Inc. & Global Blood Therapeutics, Inc.*, Case  
23 No. 160101/2025 (N.Y. Sup. Ct. – N.Y. Cnty.). Status: Plaintiff filed the  
24 complaint in the Supreme Court of the State of New York, New York County  
25 on July 31, 2025. Plaintiff served the complaint on Defendant Pfizer Inc. on  
26 July 31, 2025, and on Defendant Global Blood Therapeutics, Inc. on August  
27 4, 2025.
- 28

1 13. *Hahssan Cheaver v. Pfizer Inc. & Global Blood Therapeutics, Inc.*, Case No.  
2 816046/2025E (N.Y. Sup. Ct. – Bronx Cnty.). Status: Plaintiff filed the  
3 complaint in the Supreme Court of the State of New York, New York County  
4 on July 30, 2025. Plaintiff served the Complaint on Defendant Pfizer Inc. on  
5 July 30, 2025, and on Defendant Global Blood Therapeutics, Inc. on August  
6 4, 2025.

7 14. *Egnonnoumimon Houecande v. Pfizer Inc. & Global Blood Therapeutics,*  
8 *Inc.*, Case No. 816045/2025E (N.Y. Sup. Ct. – Bronx Cnty.). Status: Plaintiff  
9 filed the complaint in the Supreme Court of the State of New York, New York  
10 County on July 30, 2025. Plaintiff served the complaint on Defendant Pfizer  
11 Inc. on July 30, 2025, and on Defendant Global Blood Therapeutics, Inc. on  
12 August 4, 2025.

13 One case previously pending in California state court—*Leona Smith v. Global Blood*  
14 *Therapeutics, Inc. & Pfizer Inc.*, Case No. 24-CIV-08190 (Cal. Super. Ct. – San Mateo Cnty.)—has  
15 been dismissed. Plaintiff filed a request for voluntary dismissal on June 30, 2025.

16 **11. Relief**

17 **a. Plaintiff’s Statement**

18 Plaintiff seeks a jury trial and the following categories of damages: past, present and future  
19 general damages in an amount to be determined at trial; For past, present and future special damages,  
20 including but not limited to past, present and future lost earnings, economic damages and others, in  
21 an amount to be determined at trial; any appropriate punitive or exemplary damages; any appropriate  
22 statutory damages; for costs of suit; for interest as allowed by law; for attorney’s fees and costs as  
23 applicable; for treble damages as applicable; for such other and further relief as the court may deem  
24 proper.

25 **b. Defendants’ Statement**

26 Defendants dispute that they are liable to Plaintiff for any damages or other relief. If liability  
27 is established, damages expert(s) would likely be required to calculate damages, if any. Defendants  
28

1 have not yet filed their Answer but expect to do so, if appropriate, following the resolution of their  
2 forthcoming Motion to Dismiss. Defendants reserve all rights to seek all appropriate relief.

3 **12. Settlement and ADR**

4 The Parties have engaged in informal discussions and exchanged information in an effort to  
5 resolve this matter and those discussions are ongoing. Mediation is also scheduled to take place on  
6 September 9, 2025.

7 **13. Other References**

8 The Parties agree that this case is not suitable for reference to a special master or the Judicial  
9 Panel on Multidistrict Litigation.

10 **14. Narrowing Issues**

11 The Parties have not agreed on any issues that can be narrowed at this time.

12 **15. Expedited Trial Procedure**

13 The Parties agree that this case is not suitable for the Expedited Trial Procedure set forth in  
14 General Order 64, Attachment A.

15 **16. Scheduling**

16 The Court entered a Case Management and Scheduling Order on February 14, 2025 (Dkt.  
17 32), attached hereto as *Exhibit A*.

18 **17. Trial**

19 The Court has sent this case for a jury trial to commence on June 7, 2027 and last 12-15 days.

20 **18. Disclosure of Non-Party Interested Entities or Persons**

21 Plaintiff will file his Certificate of Interested Parties. Plaintiff does not have conflicts or  
22 interests to report outside of the parties.

23 Defendants filed their Certificate of Interested Entities or Persons on January 24, 2025. As  
24 disclosed therein, Pfizer Inc. is a publicly held corporation and there is no parent corporation or  
25 publicly held corporation that owns 10% or more of its common stock. Global Blood Therapeutics,  
26 Inc. is a wholly-owned subsidiary of Pfizer. Other than the parties, there is no other conflict or  
27 interest to report (Dkt. 21).

28 **19. Professional Conduct**

1 All attorneys of record for the Parties have reviewed the Guidelines for Professional Conduct  
2 for the Northern District of California.

3 **20. Other**

4 At this time, the Parties are not aware of other matters that may facilitate the resolution of  
5 this matter.

6  
7 DATED: August 14, 2025

8 By: /s/ Caitlyn Prichard Miller  
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**SIGNATURE ATTESTATION**

I, Teresa M. Wogoman, am the ECF User whose ID and password are being used to file this document. In compliance with Civil Local Rule 5-1, I hereby attest that all counsel whose e-signatures (/s/) appear on this document concurred in this filing.

DATED: August 14, 2025

By: /s/ Teresa M. Wogoman

# **EXHIBIT A**

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United States District Court  
Northern District of California

UNITED STATES DISTRICT COURT  
NORTHERN DISTRICT OF CALIFORNIA

Tirrell Allen  
  
Plaintiff,  
  
v.  
Global Blood Therapeutics, Inc, Pfizer, Inc.,  
et al  
  
Defendant.

Case No. [24-cv-07786-TLT](#)

**CASE MANAGEMENT AND  
SCHEDULING ORDER**

Re: Dkt. No. 29

Pursuant to Federal Rule of Civil Procedure 16 and Civil Local Rule 16-10, **THE  
FOLLOWING DEADLINES ARE HEREBY ORDERED:**

1. TRIAL DATE: June 7, 2027  
No. of Days: 12–15 days  
Courtroom 9, 19th Floor  
Jury Trial
2. FINAL PRETRIAL CONFERENCE: May 6, 2027, at 3:00 p.m. [in person]  
**LEAD COUNSEL WHO WILL TRY  
THE CASE MUST ATTEND**  
Joint Pretrial Statement (including  
objections, motions in limine and jury  
instructions): April 8, 2027
3. DISPOSITIVE MOTIONS AND  
DAUBERT MOTIONS: Last day to *file* dispositive motion and  
*Daubert* motions: December 11, 2026  
  
Last day to be heard: February 2, 2027,  
2:00 p.m. [in person]  
See Civil Local Rules for notice and  
filing requirements.
4. FACT DISCOVERY CUT-OFF: December 15, 2025
5. EXPERT REPORTS: Opening reports by January 20, 2026  
Rebuttal reports by March 13, 2026
6. EXPERT DISCOVERY CUT-OFF April 10, 2026

United States District Court  
Northern District of California

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- 7. ADR: Settlement conference with Private Mediator to be completed by August 22, 2025.
- The parties are ordered to provide the Court with the name of the mediator and the mediation schedule no later than March 7, 2025.
- 8. LAST DAY TO AMEND PLEADINGS May 22, 2025
- 9. JOINT STATEMENT RE. RELATED CASES IN FEDERAL AND STATE COURT INCLUDING THE POSTURE OF EACH CASE. March 7, 2025
- 10. FURTHER STATUS CONFERENCE: July 10, 2025, 2:00 p.m.; December 11, 2025, 2:00 p.m.

**IT IS FURTHER ORDERED** that parties and counsel refer to and comply with Judge Thompson’s Civil Standing Order and Civil Pretrial and Jury Trial Standing Order or Civil Pretrial and Bench Trial Standing Order located on the court’s website (<https://cand.uscourts.gov/trina-l-thompson/>).

**JURY TRIAL**

The Court will take cause challenges and discuss hardship claims at side bar. The Court will inform counsel which hardship claims and cause challenges will be granted but will not announce those dismissals until the selection process is completed. Peremptory challenges will be made in writing. The Court will strike at one time those with meritorious hardship claims, those excused for cause, and those challenged peremptorily, and then seat the first six to eight people remaining in numerical order.

The Court will send out to prospective jurors in advance of trial an electronic questionnaire soliciting information. In addition, if the parties wish to submit proposed voir dire questions, the parties must meet and confer and file a **joint set** of proposed questions that the Court may add to the electronic questionnaire. The parties will receive the responses prior to the in-court voir dire.

United States District Court  
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**SCHEDULING ORDER MODIFICATIONS**

No provision of this order may be changed except by written order of this Court upon its own motion or upon motion of one or more parties made pursuant to Civil Local Rule 7-11 with a showing of good cause. Parties may file a formal brief, but a letter brief will suffice. The requesting party shall serve the opposing party on the same day the motion is filed and the opposing party shall submit a response as soon as possible but no later than four days after service.

If the modification sought is an extension of a deadline contained herein, the motion must be brought at least seven (7) days before expiration of that deadline. **The parties may not modify the pretrial schedule by stipulation.** A conflict with a court date set after the date of this order does not constitute good cause. The parties are advised that if they stipulate to a change in the discovery schedule, they do so at their own risk. The only discovery schedule that the Court will enforce is the one set in this order.

**IT IS SO ORDERED.**

Dated: February 13, 2025

  
TRINA L. THOMPSON  
United States District Judge

November 7, 2027