

FDA FREEDOM OF INFORMATION ACT APPEAL

SUBMITTED VIA EMAIL

August 29, 2022

Director, Office of the Executive Secretariat
US Food & Drug Administration
5630 Fishers Lane, Room 1050
Rockville, MD 20857
E-mail: FDAFOIA@fda.hhs.gov

Re: Appeal of FDA Control # 2022-4857 (IR#0802O)

Dear Sir or Madam:

This firm represents Informed Consent Action Network (“**ICAN**”). On behalf of ICAN, on June 30, 2022, we submitted a request for records from the files of Food and Drug Administration (“**FDA**”) pursuant to the Freedom of Information Act (5 U.S.C. § 552, as amended) (“**FOIA**”). The FDA designated the request as FOIA Control # 2022-4857 (the “**FOIA Request**”). On August 25, 2022, the FDA issued a final response to the FOIA Request (the “**Final Response**”). ICAN writes now to appeal the Final Response.

FOIA Control # 2022-4857 (IR#0802O)

On June 30, 2022, ICAN submitted a request to the FDA for the following documents:

All “reports of possible concern based on the data mining results” the FDA shared with the CDC pursuant to section 2.5 of the VAERS Standard Operating Procedures for COVID-19. [See <https://www.cdc.gov/vaccinesafety/pdf/VAERS-v2-SOP.pdf>]

Information helpful to fulfilling the request: The FDA’s Center for Biologics Evaluation and Research is the likely custodian of responsive records.

(Exhibit 1.)¹

¹ All “Exhibits” referenced herein are appended to this letter.

On July 5, 2022, FDA acknowledged the FOIA request and assigned it FOIA Control # 2022-4857. (Exhibit 2.)

A. FDA's Final Response

On August 25, 2022, FDA issued a final response letter. The letter stated in part,

We are denying your entire request. Specifically, we are denying data mining reports.

The following exemption(s) of FOIA, 5 U.S.C. 552, is the authority for denying you access to the non-disclosable material: Exemption (b)5 Certain interagency and intra-agency communications.

(Exhibit 3.)

B. Argument

FDA improperly withheld documents and information pursuant to FOIA Exemption 5 and failed to conduct an adequate search. For the reasons set forth below, ICAN appeals FDA's Final Response.

1. The FDA Improperly Withheld Records Under FOIA Exemption 5

FDA has not properly demonstrated that the withheld records fall under the scope of Exemption 5. "An agency withholding responsive documents from a [FOIA] release bears the burden of proving the applicability of the claimed exemptions." *American Civil Liberties Union v. DOD*, 628 F.3d 612, 619 (D.C. Cir. 2011). "Exemption 5 claims must be supported with specificity and detail." *Judge Rotenberg Educ. Ctr., Inc. v. United States FDA*, 376 F. Supp. 3d 47, 65 (D.D.C. 2019) (citations omitted). The document must be: (1) an inter-agency or intra-agency document; (2) "predecisional"; and (3) deliberative. *Tigue v. United States DOJ*, 312 F.3d 70, 76 (2nd Cir. 2002). Courts have defined 'predecisional' as records "prepared in order to assist an agency decision[maker] in arriving at his decision." *Nat'l Day Laborer Org. Network v. United States Immigration & Customs Enf't*, 486 F. Supp. 3d 669, 690 (S.D.N.Y. 2020). Whereas 'deliberative' has been defined by the courts as records "related to the process by which policies are formulated." *Id.* This standard requires the agency to explain (i) "the nature of the specific deliberative process involved," (ii) "the function and significance of the documents in that process," and (iii) "the nature of the decision making authority vested in the document's author and recipient." *Brennan Ctr. for Justice at NY Univ. Sch. of Law v. Dep't. of Homeland Sec.*, 331 F. Supp. 3d 74, 93-94 (S.D.N.Y. 2018).

Additionally, to carry its burden, the agency "must demonstrate that (A) the materials at issue are covered by the deliberative process privilege, and (B) it is reasonably foreseeable that release of those material *would cause harm* to an interest protected by that privilege." (emphasis added) *Reporters Comm. for Freedom of the Press v. FBI*, 3 F.4th 350, 361 (D.C. Cir. 2021) (citing *Machado Amadis v. U.S. Dep't of State*, 971 F.3d 364, 370 (D.C. Cir. 2020); 5 U.S.C. §

552(a)(8)(A)(i)(I)). “In the context of withholdings made under the deliberative process privilege, the foreseeability requirement means that agencies must concretely explain how disclosure ‘would’—not ‘could’—adversely impair internal deliberations.” *Reporters Comm. for Freedom of the Press*, 3 F.4th. at 369-70 (quoting *Machado Amadis*, 971 F.3d at 371).

Even if the deliberative process privilege applies, it “does not protect documents in their entirety; if the government can segregate and disclose non-privileged factual information within a document, it must.” *Nat’l Day Laborer Org. Network v. United States Immigration & Customs Enf’t*, 486 F. Supp. 3d 669, 689 (S.D.N.Y. 2020) (quoting *Loving v. Dep’t of Def.*, 550 F.3d 32, 38 (D.C. Cir. 2008)). “Any reasonably segregable portion of a record shall be provided to any person requesting such record after deletion of the portions which are exempt under this subsection.” 5 U.S.C. § 552(b). Only factual material that is “inextricably intertwined with exempted portions” of the documents need not be disclosed. *Johnson v. Exec. Office for U.S. Attorneys*, 310 F.3d 771, 776 (D.C. Cir. 2002). The government has the “burden of demonstrating that no reasonably segregable information exists within . . . documents withheld.” *Loving v. Dep’t of Defense*, 550 F.3d 32, 41(D.C. Cir. 2008). “[T]he ultimate objective of exemption 5 is to safeguard the deliberative process of agencies, not the paperwork generated in the course of that process.” *Nat’l Wildlife Fed’n v. U.S. Forest Serv.*, 861 F.2d 1114, 1119 (9th Cir. 1988).

There are three reasons why FDA’s application of Exemption 5 was improper. First, FDA has not provided the specificity and detail required to deny the entire FOIA Request by invoking Exemption 5. *Judge Rotenberg Educ. Ctr., Inc.*, 376 F. Supp. 3d at 65. For example, FDA’s final response does not explain specifically how the documents qualify as (1) an inter-agency or intra-agency document; (2) predecisional; and (3) deliberative. The agency did not explain the nature of the deliberative process involved, the function and significance of the documents withheld under the deliberative process, or the nature of the decision-making authority vested in the documents author and recipient. Instead of providing the specificity and detail that FOIA requires, FDA – without further explanation – only vaguely cited “Exemption (b)5”, and a few other sections of the Code of Federal Regulations in its Final Response. (See **Exhibit 3**.) Thus, the applicability of Exemption 5 has not been proven. *American Civil Liberties Union*, 628 F.3d at 619; *Tigue*, 312 F.3d at 76; *Brennan Ctr. for Justice at NY Univ. Sch. of Law*, 331 F. Supp. 3d at 93-94.

Second, FDA’s Final Response did not provide any information as to how it is reasonably foreseeable that the release of the withheld materials would cause harm by adversely impairing internal deliberations. *Reporters Comm. for Freedom of the Press*, 3 F.4th. at 369-70.

Lastly, the FDA has not provided any details as to why factual information does not exist or why any factual information within the withheld documents is not reasonably segregable. *Loving*, 550 F.3d at 41. The information requested concerns data mining results. Results, by their nature, are conclusions and final determinations which are not pre-decisional. Any final conclusion or final documents should be produced. In addition, there is a high likelihood that the withheld documents contain factual information. Moreover, the reports themselves likely have dates of review and completion, attachments, and other purely factual information that are reasonably segregable from possibly exempt portions. The agency should only utilize Exemption

5 to safeguard the deliberative process, not all the paperwork generated in the course of that process. *Nat'l Wildlife*, 861 F.2d at 1119.

FDA has provided no detail about the segregability of the withheld records, beyond the following:

The following section of the implementing regulations of FDA and reason(s) applicable to this denial are contained in the CFR, Title 21

- **Section 20.62 Intra-agency memoranda consisting of opinions, recommendations, and policy discussions within the deliberative process of FDA, from which factual information is not reasonably segregable.**

Thus, FDA failed to demonstrate that no reasonably segregable information exists within the documents withheld. *Loving*, 550 F.3d at 41.

Therefore, for these reasons, FDA has not met its burden in proving the applicability of Exemption 5, and the withheld records should be immediately released. *American Civil Liberties Union*, 628 F.3d at 619.

2. FDA Failed to Conduct an Adequate Search

FDA has failed to conduct an adequate search of the requested records. An agency's search is adequate only if it is "reasonably calculated to uncover all relevant documents." *Zemansky v. E.P.A.*, 767 F.2d 569, 571 (9th Cir. 1985) (quoting *Weisberg v. U.S. Dep't. of Justice*, 745 F.2d 1476, 1485 (D.C. Cir. 1984)) (internal quotation marks omitted). "An agency fulfills its obligations under FOIA if it can demonstrate *beyond material doubt* that its search was reasonably calculated to uncover all relevant documents." *Def. of Wildlife v. United States Border Patrol*, 623 F. Supp. 2d 83, 91 (D.D.C. 2009) (quoting *Valencia-Lucena v. U.S. Coast Guard*, 180 F.3d 321, 325 (D.C. Cir. 1999)) (emphasis added). To satisfy its FOIA obligations, an agency needs to adequately describe the scope and methods of its searches, which can reasonably be expected to uncover the records sought and demonstrate that the places most likely to contain responsive materials were searched. *Davidson v. E.P.A.*, 121 F. Supp. 2d 38, 39 (D.D.C. 2000). At minimum, the agency must specify "what records were searched, by whom, and through what process." *Steinberg v. U.S. Dep't. of Justice*, 23 F.3d 548, 552 (D.C. Cir. 1994).

FDA search was inadequate because it failed to specify what records were searched, by whom, and through what process. *Steinberg*, 23 F.3d 552. Therefore, FDA did not fulfill its obligations under FOIA of demonstrating beyond material doubt that its search was reasonably calculated to uncover all relevant documents. *Valencia-Lucena*, 180 F.3d at 325.

C. Appellate Request

Given the foregoing, ICAN hereby appeals and requests that the documents responsive to the FOIA Requests be produced within 20 days of this appeal. Thank you for your time and

attention to this matter. If you require any additional information, please contact us at **(212) 532-1091** or through email at **foia@sirillp.com**.

Very truly yours,

/s/ Aaron Siri

Aaron Siri, Esq.

Elizabeth A. Brehm, Esq.

Colin Farnsworth, Esq.

Enclosures

Exhibit 1

FDA FREEDOM OF INFORMATION ACT REQUEST

VIA ONLINE PORTAL

June 30, 2022

Food and Drug Administration
Division of Freedom of Information
Office of the Secretariat, OC
5630 Fishers Lane, Room 1035
Rockville, MD 20857

Re: "Reports of Possible Concern" – VAERS Standard Operating Procedures for COVID-19 (IR#08020)

Dear Sir or Madam:

This firm represents the Informed Consent Action Network ("ICAN"). On behalf of ICAN, please provide the following records to foia@sirillp.com in electronic form:

All "reports of possible concern based on the data mining results" the FDA shared with the CDC pursuant to section 2.5 of the VAERS Standard Operating Procedures for COVID-19.¹

Information helpful to fulfilling the request: The FDA's Center for Biologics Evaluation and Research is the likely custodian of responsive records.

We ask that you waive any and all fees or charges pursuant to 5 U.S.C. § 552(a)(4)(A)(iii). ICAN is a not-for-profit news media organization whose mission is to raise public awareness about vaccine safety and to provide the public with information to give informed consent. (**Exhibit A.**) As part of its mission, ICAN actively investigates and disseminates information regarding vaccine safety issues for free, including through its website,² a weekly health news and talk show,³ and through press events and releases. ICAN is seeking the information in this FOIA request to allow it to contribute to the public understanding of the government's vaccine safety programs, including the government's efforts to promote vaccine safety. The information ICAN is requesting will not contribute to any commercial activities. Therefore, ICAN should be properly categorized as a

¹ See <https://www.cdc.gov/vaccinesafety/pdf/VAERS-v2-SOP.pdf>.

² <https://www.icandecide.org/>.

³ <https://thehighwire.com/>.

media requester, and it is entitled to the search and processing privileges associated with such a category designation. Accordingly, ICAN will be forced to challenge any agency decision that categorizes it as any other category of requester.

Please note that the FOIA provides that if only portions of a requested file are exempted from release, the remainder must still be released. We therefore request that we be provided with all non-exempt portions which are reasonably segregable. We further request that you describe any deleted or withheld material in detail and specify the statutory basis for the denial as well as your reasons for believing that the alleged statutory justification applies. Please also separately state your reasons for not invoking your discretionary powers to release the requested documents in the public interest. Such statements may help to avoid unnecessary appeal and litigation. ICAN reserves all rights to appeal the withholding or deletion of any information.

Access to the requested records should be granted within twenty (20) business days from the date of your receipt of this letter. Failure to respond in a timely manner shall be viewed as a denial of this request and ICAN may immediately take further administrative or legal action.

Furthermore, we specifically request that the agency provide us with an estimated date of completion for this request.

If you would like to discuss our request or any issues raised in this letter, please feel free to contact us at (212) 532-1091 or foia@sirillp.com during normal business hours. Thank you for your time and attention to this matter.

Very truly yours,

/s/ Aaron Siri

Aaron Siri, Esq.

Elizabeth A. Brehm, Esq.

Colin M. Farnsworth Esq.

Exhibit A

DECLARATION OF CATHARINE LAYTON

STATE OF TEXAS

COUNTY OF Hays

I, Catharine Layton, being duly sworn on oath do say:

1. I am the Chief Operating Officer of the Informed Consent Action Network (ICAN), a not-for-profit 501(c)(3) organization whose mission is to disseminate scientific health information to the public.

2. I have been an officer of ICAN since its founding in 2016. I oversee all day-to-day operations of the organization and all ICAN's programs. Together with our CEO and Board, I ensure that all efforts are focused on our mission statement and ensure that ICAN stays in compliance with all required rules and regulations.

3. In pursuit of its mission, ICAN relies primarily on its own investigative reporting. ICAN is both instrumental in orchestrating cutting edge investigations into the safety of various medical products, as well as widely disseminating its findings through various media channels. Most notably, ICAN's popular website hosts the organization's largest education program, The HighWire with Del Bigtree. Utilizing its media teams' 40+ years of experience in TV production and investigative journalism, The HighWire provides hours of new video content to the public each week for free.

4. The HighWire website has approximately 3.4 million weekly visitors. On Twitter, The HighWire has approximately 140,000 followers and 1 to 2.5 million impressions in a 28-day period. Between Rumble and Bitchute, The HighWire has approximately 60,000 followers and growing. Additionally, ICAN has 29,000 text subscribers and 194,245 email subscribers.

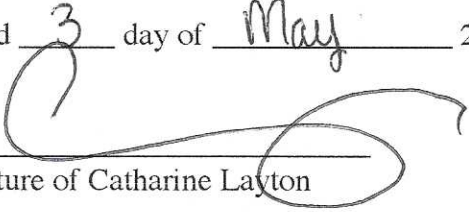
5. The size of ICAN's audience and subscribers continues to grow and is illustrative of the wide public interest in the subject of health and medical safety. Moreover, critical to ICAN's mission is its proven ability to find and review critical scientific and governmental records and meaningfully report about their social impacts.

6. One of the tools ICAN uses to gather the raw material it uses in its popular investigative reporting is the Freedom of Information Act (FOIA).

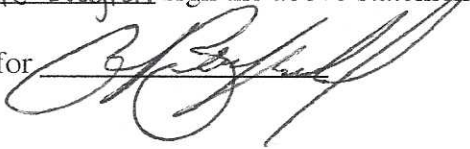
7. ICAN uses records it obtains from its FOIA requests to carry out its public mission and support its role as a non-profit news-media organization in the field of health and medical safety, but as a non-profit, ICAN does not have a commercial interest in the records it seeks through FOIA.

8. Based on what I know as the Chief Operating Officer, as well what has been demonstrated by ICAN's past and current investigative reporting, for purposes of FOIA's Fee Waiver provisions, ICAN certainly qualifies as a "representative of the news media."

Signed 3 day of May 2022


Signature of Catharine Layton

I, Amy Blackwell Notary public for the state of Texas witnessed
said Catharine Layton sign the above statement this 3 day of May, 2022
(month)

Notary Public for 

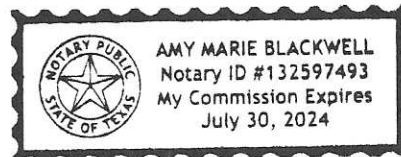


Exhibit 2



July 05, 2022

SIRI & GLIMSTAD LLP
AARON SIRI
200 PARK AVE STE 3300
NEW YORK NY 10166

In Reply refer to
FOIA Control #:
2022-4857

Requester reference:
IR#08020

Dear Requester:

The Food and Drug Administration (FDA) has received your Freedom of Information Act (FOIA) request for records regarding:

All "reports of possible concern based on the data mining results" the FDA shared with the CDC pursuant to section 2.5 of the VAERS Standard Operating Procedures for COVID-19.

We will respond as soon as possible and may charge you a fee for processing your request. If your informational needs change, and you no longer need the requested records, please contact us to cancel your request, as charges may be incurred once processing of your request has begun. For more information on processing fees, please see <http://www.fda.gov/RegulatoryInformation/FOI/FOIAFees/default.htm>.

Due to an increase in the number of incoming requests, we may be unable to comply with the twenty-working-day time limit in this case, as well as the ten additional days provided by the FOIA. The actual processing time will depend on the complexity of your request and whether sensitive records, voluminous records, extensive search, and/or consultation with other HHS components or other executive branch agencies are involved. Please note that requests for medical device approval records (e.g. 510K, PMA, DEN) may take up to 18 to 24 months to process.

If you have any questions about your request, please call Wilson M. Russ, Freedom Of Information Specialist, at (301) 796-8981 or write to us at:
Food and Drug Administration
Division of Freedom of Information
5630 Fishers Lane, Room 1035
Rockville, MD 20857

If you call or write, use the FOIA control number provided above which will help us to answer your questions more quickly.

You also have the right to seek dispute resolution services from:

Office of Government Information Services
National Archives and Administration
8601 Adelphi Road – OGIS
College Park, MD 20740-6001
Telephone: 202-741-5770
Toll-Free: 1-877-684-6448
Email: ogis@nara.gov
Fax: 202-741-5769

and/or

FDA FOIA Public Liaison
Office of the Executive Secretariat
US Food Administration
5630 Fishers Lane, Room 1050
Email: FDAFOIA@fda.hhs.gov

Sincerely,

SARAH KOTLER
Director

Exhibit 3

Annalise Beube

From: Kotler, Sarah <Sarah.Kotler@fda.hhs.gov>
Sent: Thursday, August 25, 2022 5:23 AM
To: S&G Information Request Staff
Subject: FDA FOIA 2022-4857

Follow Up Flag: Follow up
Flag Status: Flagged

Dear Requester:

The Food and Drug Administration (FDA) has completed processing your request for records under the Freedom of Information Act (FOIA).

We are denying your entire request. Specifically, we are denying data mining reports.

The following exemption(s) of FOIA, 5 U.S.C. 552, is the authority for denying you access to the non-disclosable material: Exemption (b)5 Certain interagency and intra-agency communications. We have included citations to the FOIA and FDA's regulations for your information.

Section 5.31 (e) of the implementing regulations of the Department of Health and Human Services (DHHS) is applicable to this denial. The regulations are contained in the Code of Federal Regulations (CFR), Title 45.

The following sections of the implementing regulations of FDA and reason(s) applicable to this denial are contained in the CFR, Title 21

- Section 20.62 Intra-agency memoranda consisting of opinions, recommendations, and policy discussions within the deliberative process of FDA, from which factual information is not reasonably segregable.

FDA's Regulations at CFR Part 20 are available at:

http://www.access.gpo.gov/nara/cfr/waisidx_04/21cfr20_04.html

You have the right to appeal this determination. By filing an appeal, you preserve your rights under FOIA and give the agency a chance to review and reconsider your request and the agency's decision. Your appeal must be mailed within 90 days from the date of this response, to: Director, Office of the Executive Secretariat, US Food & Drug Administration, 5630 Fishers Lane, Room 1050, Rockville, MD 20857, E-mail: FDAFOIA@fda.hhs.gov. Please clearly mark both the envelope and your letter "FDA Freedom of Information Act Appeal."

If you would like to discuss our response before filing an appeal to attempt to resolve your dispute without going through the appeals process, please contact Sarah Kotler at 301-796-8976. You may also contact the FDA FOIA Public Liaison for assistance at: Office of the Executive Secretariat, US Food & Drug Administration, 5630 Fishers Lane, Room 1050, Rockville, MD 20857, E-mail: FDAFOIA@fda.hhs.gov.

If you are unable to resolve your FOIA dispute through our FOIA Public Liaison, the Office of Government Information Services (OGIS), the Federal FOIA Ombudsman's office, offers mediation services to help resolve disputes between FOIA requesters and Federal agencies. The contact information for OGIS is as follows: Office of Government Information Services, National Archives and Records Administration, 8601 Adelphi Road—OGIS, College Park, MD 20740-6001; telephone at 202-741-5770; toll free at 1-877-684-6448; or facsimile at 202-741-5769; e-mail at ogis@nara.gov.

Sincerely yours,

Sarah Kotler
Director
Division of Freedom of

Information

Sarah B. Kotler, J.D.
Director, Division of Freedom of Information
US FDA
301-796-8976