



Christine, an unincorporated woman <cmssyc@gmail.com>

FDA Receipt of FOI Request Control # 2024-7837

FDA_FOI@fda.gov <FDA_FOI@fda.gov>
To: cmssyc@gmail.com
Cc: FDAFOIA@fda.hhs.gov

Wed, Sep 4, 2024 at 8:08 AM

Control number: 2024-7837

Please find the attached acknowledgement regarding your FOIA request.

Note: Do NOT reply directly to this E-mail

 **Acknowledgement Letter.PDF**
62K



September 04, 2024

FLUORIDE FREE PEEL
CHRISTINE: HOUSE OF MASSEY



In Reply refer to
FOIA Control #:
2024-7837

Requester reference:

Dear Requester:

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

All studies in the possession/custody/control of the FDA, authored by anyone, anywhere: 1. - that scientifically prove/provide evidence of the existence of any alleged "SARS-COV-2" (including "variants") (showing at minimum that the alleged particles with a specific "genome" and proteins exist and cause the illness/symptoms that they are alleged to cause) Note: Scientific proof/evidence is not opinions, speculation, declarations, review papers or descriptive studies. Scientific evidence requires use of the scientific method to test falsifiable hypotheses through valid, rigorous, repeatable controlled experiments. 2. - or, that even describe the purification of particles that are alleged to be "SARS-COV-2" directly from bodily fluid/tissue/excrement of so-called "hosts" (without adding any sources of genetic material or proteins) with purification confirmed via EM images (the images must be available as well) Purify = separate from everything else in the clinical sample. I am aware that according to "virus" dogma a "virus" requires host cells in order to replicate and am not seeking records describing the replication of a "virus" without "host" cells, or that describe a suspected "virus" floating in a vacuum or a strict fulfillment of Koch's Postulates. I am simply seeking records that describe purification (separation from everything else in the "host" sample). I am not seeking private patient records. 3. - or, wherein the purported "genome" of any alleged "SARS-COV-2" was found intact in the bodily fluid/tissue/excrement of a so-called "host" (as opposed to fabricated/assembled in silico, aka a computer model) 4. - or, that scientifically demonstrate the claimed contagious nature of the illness / symptoms that are allegedly caused by purported "SARS-COV-2". Send via email. I do not agree to this order or "personal information" being shared with 3rd parties including those that provide FOIA services, without my explicit written consent.

In processing your FOIA request, FDA will apply, as appropriate, the FOIA exemptions in 5 USC 552(b) and the foreseeable harm standard in 5 USC 552(a)(8)(i). 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>. PLEASE NOTE: HOURLY RATES FOR SEARCH AND REVIEW INCREASED FOR ALL REQUESTS RECEIVED ON OR AFTER JUNE 1, 2023.

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 Sarah B. Kotler, Director, Division Of Freedom Of Information, at (301) 796-8976 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

and/or

FDA FOIA Public Liaison
Office of the Executive Secretariat
US Food and Drug Administration

College Park, MD 20740-6001
Telephone: 202-741-5770
Toll-Free: 1-877-684-6448
Email: ogis@nara.gov
Fax: 202-741-5769

5630 Fishers Lane, Room 1050
Rockville, MD 20857
Email: FDAFOIA@fda.hhs.gov

Sincerely,

SARAH KOTLER
Director



Christine, an unincorporated woman <cmssyc@gmail.com>

FDA Notification of FOI Expedite Request Denied Control # 2024-7837

FDA_FOI@fda.gov <FDA_FOI@fda.gov>
To: cmssyc@gmail.com
Cc: FDAFOIA@fda.hhs.gov

Tue, Sep 10, 2024 at 5:08 PM

Control number: 2024-7837

Please find the attached response regarding your expedite request.

Note: Do NOT reply directly to this E-mail

 **Expedited Processing Denied.PDF**
55K



September 10, 2024

FLUORIDE FREE PEEL
CHRISTINE: HOUSE OF MASSEY



In Reply refer to
FOIA Control #:
2024-7837

Requester reference:

Dear Requester:

This is in reference to your request(s) for record(s) from the Food and Drug Administration (FDA) pursuant to the Freedom of Information Act (FOIA).

All studies in the possession/custody/control of the FDA, authored by anyone, anywhere: 1. - that scientifically prove/provide evidence of the existence of any alleged "SARS-COV-2" (including "variants") (showing at minimum that the alleged particles with a specific "genome" and proteins exist and cause the illness/symptoms that they are alleged to cause) Note: Scientific proof/evidence is not opinions, speculation, declarations, review papers or descriptive studies. Scientific evidence requires use of the scientific method to test falsifiable hypotheses through valid, rigorous, repeatable controlled experiments. 2. - or, that even describe the purification of particles that are alleged to be "SARS-COV-2" directly from bodily fluid/tissue/excrement of so-called "hosts" (without adding any sources of genetic material or proteins) with purification confirmed via EM images (the images must be available as well) Purify = separate from everything else in the clinical sample. I am aware that according to "virus" dogma a "virus" requires host cells in order to replicate and am not seeking records describing the replication of a "virus" without "host" cells, or that describe a suspected "virus" floating in a vacuum or a strict fulfillment of Koch's Postulates. I am simply seeking records that describe purification (separation from everything else in the "host" sample). I am not seeking private patient records. 3. - or, wherein the purported "genome" of any alleged "SARS-COV-2" was found intact in the bodily fluid/tissue/excrement of a so-called "host" (as opposed to fabricated/assembled in silico, aka a computer model) 4. - or, that scientifically demonstrate the claimed contagious nature of the illness / symptoms that are allegedly caused by purported "SARS-COV-2". Send via email. I do not agree to this order or "personal information" being shared with 3rd parties including those that provide FOIA services, without my explicit written consent.

The Electronic Freedom of Information Act (EFOIA) Amendments of 1996 amended the FOIA by adding section (a)(6)(E), 5 U.S.C. 552(a)(6)(E), to require agencies to consider requests for expedited processing and grant them whenever a "compelling need" is shown and in other cases as determined by the agency. The term "compelling need" is defined as (1) involving "an imminent threat to the life or physical safety of an individual," or (2) in the case of a request made by "a person primarily engaged in disseminating information, urgency to inform the public concerning actual or alleged Federal Government activity."

I have determined that your request for expedited processing does not meet the criteria under the FOIA. You have not demonstrated a compelling need that involves an imminent threat to the life or physical safety of an individual. Neither have you demonstrated that there exists an urgency to inform the public concerning actual or alleged Federal Government activity. Therefore, I am denying your request for expedited processing. The responding agency office will process your request in the order in which it was received.

In accordance with 45 CFR § 5.61 and 21 CFR § 20.41(b)(5), you have the right to appeal this determination. Your appeal should clearly identify the agency determination that is being appealed. It would be helpful if you provide specific reasons explaining why you believe the agency's adverse determination should be reconsidered. 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, U.S. Food & Drug Administration, 5630 Fishers Lane, Room 1050, Rockville, MD 20857, or emailed within 90 days from the date of this response to FDAFOIA@fda.hhs.gov. Please clearly mark both the envelope and your letter "FDA Freedom of Information Act Appeal." Items arriving or delivered after 5 p.m. Eastern Time will be deemed received on the next workday.

You may also contact the FDA FOIA Public Liaison, Office of the Executive Secretariat, 5630 Fishers Lane, Room 1050, Rockville, MD 20857; email: 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: Office of Government Information

Services, National Archives and Records Administration, 8601 Adelphi Road—OGIS, College Park, MD 20740-6001,
Telephone: 202-741-5770, Toll-Free: 1-877-684-6448, E-mail: ogis@nara.gov, Fax: 202-741-5769.

Sincerely,

SARAH KOTLER
Director



Christine, an unincorporated woman <cmssyc@gmail.com>

FDA Notification of FOI Expedite Request Denied Control # 2024-7837

Christine, an unincorporated woman <cmssyc@gmail.com>
To: FDA_FOI@fda.gov, "Kotler, Sarah" <sarah.kotler@fda.hhs.gov>
Cc: FDAFOIA@fda.hhs.gov

Wed, Sep 11, 2024 at 10:17 AM

Notice to agent is notice to principal; notice to principal is notice to agent.

Good morning SARAH KOTLER,

Thank you for your letter. I am disappointed by your decision and it suggests a cavalier attitude towards all the loss, harm and suffering that results from pseudoscientific claims of "viruses".

Capitalizing "FLUORIDE FREE PEEL" and "CHRISTINE: HOUSE OF MASSEY" is incorrect. I ask that you kindly send a corrected letter.

Regards,
Christine
[Quoted text hidden]



Christine, an unincorporated woman <cmssyc@gmail.com>

FDA FOIA 2024-7837

Kotler, Sarah <Sarah.Kotler@fda.hhs.gov>
To: "Christine, an unincorporated woman" <cmssyc@gmail.com>

Fri, Sep 13, 2024 at 4:37 PM

Please find attached the response to your request.

Sincerely,

Sarah B. Kotler, J.D.

Director, Division of Freedom of Information

US FDA

301-796-8976

 **2024-7837.pdf**
208K



September 13, 2024

By email to: cmssyc@gmail.com

Dear Requester,

I am writing in response to your request for:

All studies in the possession/custody/control of the FDA, authored by anyone, anywhere: 1. - that scientifically prove/provide evidence of the existence of any alleged "SARS-COV-2" (including "variants") (showing at minimum that the alleged particles with a specific "genome" and proteins exist and cause the illness/symptoms that they are alleged to cause) Note: Scientific proof/evidence is not opinions, speculation, declarations, review papers or descriptive studies. Scientific evidence requires use of the scientific method to test falsifiable hypotheses through valid, rigorous, repeatable controlled experiments. 2. - or, that even describe the purification of particles that are alleged to be "SARS-COV-2" directly from bodily fluid/tissue/excrement of so-called "hosts" (without adding any sources of genetic material or proteins) with purification confirmed via EM images (the images must be available as well) Purify = separate from everything else in the clinical sample. I am aware that according to "virus" dogma a "virus" requires host cells in order to replicate and am not seeking records describing the replication of a "virus" without "host" cells, or that describe a suspected "virus" floating in a vacuum or a strict fulfillment of Koch's Postulates. I am simply seeking records that describe purification (separation from everything else in the "host" sample). I am not seeking private patient records. 3. - or, wherein the purported "genome" of any alleged "SARS-COV-2" was found intact in the bodily fluid/tissue/excrement of a so-called "host" (as opposed to fabricated/assembled in silico, aka a computer model) 4. - or, that scientifically demonstrate the claimed contagious nature of the illness / symptoms that are allegedly caused by purported "SARS-COV-2". Send via email. I do not agree to this order or "personal information" being shared with 3rd parties including those that provide FOIA services, without my explicit written consent.

The FDA does not regulate or treat viruses. The FDA is responsible for protecting the public health by ensuring the safety, efficacy, and security of human and veterinary drugs, biological products, and medical devices; and by ensuring the safety of our nation's food supply, cosmetics, and products that emit radiation. For general information on FDA's involvement with SARS-COV-2, please see: [Coronavirus Disease 2019 \(COVID-19\) | FDA](#).

In accordance with 45 CFR § 5.61 and 21 CFR § 20.41(b)(5), you have the right to appeal this determination. Your appeal should clearly identify the agency determination that is being appealed. It would be helpful if you provide specific reasons explaining why you believe the agency's adverse determination should be reconsidered. 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,


U.S. Food & Drug Administration, 5630 Fishers Lane, Room 1050, Rockville, MD 20857, or emailed within 90 days from the date of this response to FDAFOIA@fda.hhs.gov. Please clearly mark both the envelope and your letter “FDA Freedom of Information Act Appeal.” Items arriving or delivered after 5 p.m. Eastern Time will be deemed received on the next workday.

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 me at 301-796-8976 or Sarah.Kotler@fda.hhs.gov.

Please contact me if you have questions about your request. 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,

Sarah B. Kotler -S  Digitally signed by Sarah B. Kotler -S
Date: 2024.09.13 16:36:14 -04'00'

Sarah Kotler



Christine, an unincorporated woman <cmssyc@gmail.com>

FDA FOIA 2024-7837

Christine, an unincorporated woman <cmssyc@gmail.com>
To: "Kotler, Sarah" <Sarah.Kotler@fda.hhs.gov>

Fri, Sep 13, 2024 at 6:39 PM

Thank you Sarah, bu you forgot the key line in your latest confession letter:

...we have no responsive records.

Please send a revised letter that answers my FOIA order.

Thanks in advance,
Christine

[Quoted text hidden]