

Covid mRNA Vaccine and Pregnancy: Leading to Miscarriage or Stillbirth!

Fetal Cardiac Arrests, Fetal Pulmonary Hemorrhage, Placental Clots, Placental Failure

Theme: Science and Medicine

By Dr. William Makis

Global Research, June 16, 2023

COVID Intel

All Global Research articles can be read in 51 languages by activating the Translate Website button below the author's name.

To receive Global Research's Daily Newsletter (selected articles), click here.

Click the share button above to email/forward this article to your friends and colleagues. Follow us on <u>Instagram</u> and <u>Twitter</u> and subscribe to our <u>Telegram Channel</u>. Feel free to repost and share widely Global Research articles.

Pregnant women who took toxic, experimental COVID-19 mRNA vaccines have been suffering all kinds of complications including:

- miscarriages (fetal death <= 20wk)
- stillbirths (fetal death > 20wk)
- sudden death of mom before, during or after delivery (<u>click here</u>)
- pregnant women having heart attacks, strokes, dying in sleep (<u>click here</u>)
- sudden deaths of infants shortly after delivery (<u>click here</u>)
- injuries to infants such as myocarditis (<u>click here</u>)
- injuries to infants from breastfeeding with mRNA in milk (click here)

In this article I will look deeper into some of the identified causes of miscarriages and stillbirths (after mom took COVID-19 mRNA vaccine) such as fetal growth restriction, fetal cardiac arrest, fetal pulmonary hemorrhage, blood clots at the placenta, placental failure, etc.

I will write a separate substack on congenital malformations.

I would like to thank Substack author "WelcomeTheEagle88" for his work on finding these VAERS reports, please check out his substack, he has done some incredible work (click here).



WelcomeTheEagle88

@WELCOMETHEEAGLE88

#1 VAERS Auditor in the world https://www.vaersaware.com/ https://www.bitchute.com/channel/welcometheeagle88/

Fetus stopped growing after mRNA injection:

<u>CASE 01 (VAERS 1070770)</u> – Pregnant woman had 1st Pfizer jab on Feb.4, 2021 at 7wk5d pregnancy. Fetus stopped growing 6 days later (8wk4d), no heartbeat and she had miscarriage Feb.22, 2021 (18 days after Pfizer)

 VAERS ID:
 1070770 (history)
 Vaccinated:
 2021-02-04

 Form:
 Version 2.0
 Onset:
 2021-02-01

 Age:
 Submitted:
 0000-00-00

 Sex:
 Unknown
 Entered:
 2021-03-03

Location: Texas

Vaccination / Manufacturer Lot / Dose Site / Route COVID19: COVID19 (COVID19 (PFIZER-BIONTECH)) / PFIZER/BIONTECH | EL9269 / 1 | -/ OT

Administered by: Public Purchased by: ?

Symptoms: Foetal heart rate abnormal, Heart rate, Maternal exposure during pregnancy, Ultrasound scan SMQs:, Pregnancy, labour and delivery complications and risk factors (excl abortions and stillbirth) (narrow), Foetal disorders (narrow)

Life Threatening? No Birth Defect? No

Died? Yes

Date died: 2021-02-22 Days after onset: 21 Permanent Disability? No

Recovered? No Office Visit? No ER Visit? No ER or Doctor Vis

ER or Doctor Visit? Yes Hospitalized? No Previous Vaccinations:

Other Medications: VIT D; FOLATE; PRENATAL VITAMINS [ASCORBIC ACID;BETACAROTENE;CALCIUM SUI EATE: COLECA CIEEDOL: CYANOCORAL AMIN; EEDDOLIS : 701 OFT

SULFATE; COLECAL CIFEROL; CYANOCOBALAMIN; FERROUS; ZOLOFT

Current Illness: Preexisting Conditions:

Allergies:

Diagnostic Lab Data: Test Date: 20210203; Test Name: heartbeat; Result Unstructured Data: Test Result:152 bpm; Test Date: 20210220; Test Name: heartbeat; Result Unstructured Data: Test Result:no heartbeat; Test Date: 20210203; Test Name: ultrasound; Result Unstructured Data: Test Result:no abnormalities; Test Date: 20210220; Test Name: ultrasound; Result Unstructured Data: Test Result:fetus stopped growing

CDC Split Type: USPFIZER INC2021225027

Write-up: Maternal exposure during pregnancy; Fetus stopped growing on 09Feb21 (8w4d); no heartbeat detected; This is a spontaneous report from a contactable consumer (parent). This consumer reported information for both mother and fetus. This is a fetus report. A patient of unspecified age and gender (fetus) received first dose of BNT162B2 (PFIZER-BIONTECH COVID-19 VACCINE, lot number: EL9269), transplacental on 04Feb2021 at 14:00 at single dose for COVID-19 immunisation. The patient medical history was not reported. Concomitant medication included ergocalciferol (VIT D), folic acid (FOLATE), ascorbic acid/betacarotene/calcium sulfate/colecalciferol/cyanocobalamin/ferrous fumarate/folic acid/nicotinamide/pyridoxine hydrochloride/retinol acetate/riboflavin/thiamine mononitrate/tocopheryl acetate/zinc oxide (PRENATAL VITAMINS) and sertraline hydrochloride (ZOLOFT) at 25 mg, all transplacental. It was reported that OB exam on 03Feb21 showed healthy baby at 7weeks 5days neartbeat detected 152 bpm; no abnormalities identified via ultrasound; labs and hormone levels all within normal ranges. No issues detected. Mother received 1st dose of vaccine on 04Feb2021. Per ultrasound on 20Feb2021, fetus stopped growing on 09Feb2021 (8 weeks 4 days); no heartbeat detected. Miscarriage occurred on 22Feb2021. The fetus died on 22Feb2021. It was not reported if an autopsy was performed.; Sender"s Comments: Linked Report(s): US-PIZER INC-2021204433 same drug and reporter, different patient and event; Reported Cause(s) of Death: Fetus stopped growing on 09Feb21 (8w4d); no heartbeat detected. Miscarriage occurred 22Feb201.

<u>CASE 02 (VAERS 1340339)</u> – 35 year old Pregnant woman had 2nd Pfizer jab on April 18, 2021 at 5wk5d of pregnancy. Fetus stopped growing 5 days after Pfizer jab on April 23, 2021 and died. Miscarriage.

<u>CASE 03 (VAERS 1394084)</u> – Pregnant woman had 2nd Pfizer dose at 5.5 weeks pregnancy and fetus stopped growing immediately after her Pfizer jab. Miscarriage.

<u>CASE 04 (VAERS 2003811)</u> – 34 year old pregnant woman had 2nd Pfizer dose on Aug.5, 2021 at 2 weeks pregnancy and fetus stopped growing immediately, she suffered a miscarriage 3 weeks later on Aug.26, 2021.

<u>CASE 05 (VAERS 2070928)</u> – Pregnant woman had 2nd Pfizer jab at 3wk of pregnancy and fetus stopped growing at 4.5 to 5 weeks pregnancy resulting in spontaneous abortion.

<u>CASE 06 (VAERS 2115408)</u> – 29 year old pregnant woman had 3rd Pfizer jab at 8 weeks pregnancy and fetus stopped growing on same day as vaccination.

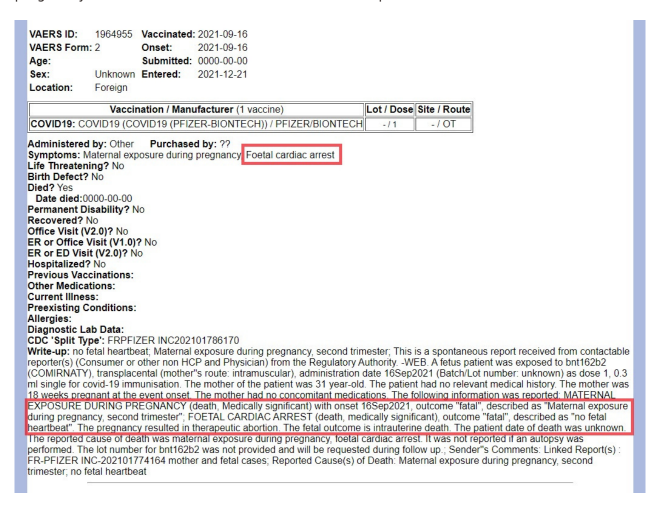
CASE 07 (VAERS 2178577) - Pregnant woman had 2nd Pfizer jab on Jul.30, 2021 and 18

days later, U/S showed intrauterine growth retardation. Medical termination of pregnancy Aug.26, 2021 (4 weeks after Pfizer jab)

<u>CASE 08 (VAERS 2180550)</u> – Pregnant woman had 3rd Pfizer jab on Dec.1, 2021 and within 2 months there was fetal growth restriction leading to fetal death on Feb.15, 2022 and medical termination of pregnancy.

Fetus had cardiac arrest:

<u>CASE 09 (VAERS 1964955)</u> – 31 year old pregnant woman had 1st Pfizer jab at 18 weeks pregnancy and suffered fetal cardiac arrest and therapeutic abortion.



<u>CASE 10 (VAERS 2009647)</u> - Pregnant woman had 1st Pfizer dose at 4wk pregnancy on May 18, 2021 and her fetus had cardiac arrest 15 days later at 6wk3d resulting in miscarriage.

VAERS ID: 2009647 Vaccinated: 2021-05-18 VAERS Form: 2 Onset: 2021-05-18 Submitted: 0000-00-00 Age: Sex: Unknown Entered: 2022-01-06 Location: Foreign Vaccination / Manufacturer (1 vaccine) Lot / Dose Site / Route COVID19: COVID19 (COVID19 (PFIZER-BIONTECH)) / PFIZER/BIONTECH ET8885 1 Administered by: Other Purchased by: ?? Symptoms: Foetal death, Scan, Foetal growth restriction, Maternal exposure during pregnancy, Foetal cardiac arrest Life Threatening? No Birth Defect? Yes Died? Yes Date died:0000-00-00 Permanent Disability? No Recovered? No Office Visit (V2.0)? No ER or Office Visit (V1.0)? No ER or ED Visit (V2.0)? No Hospitalized? No Previous Vaccinations: Other Medications: Current Illness: Preexisting Conditions: Allergies: Diagnostic Lab Data: Test Name: scans; Result Unstructured Data: Test Result: fetus stopped growing; Comments: scan at 9 weeks showed fetus stopped growing at 6w 3D CDC 'Split Type': GBPFIZER INC202101873814 Write-up scan at 9 weeks showed fetus stopped growing at 6w 3D; Heart stopped at 6weeks 3 days; Early miscarriage Maternal exposure during pregnancy, first trimester; This is a spontaneous report received from a contactable reporter(s) (Consumer or other non HCP) from the Regulatory Agency (RA). A fetus patient was exposed to bnt162b2 (BNT162B2), transplacental, administration date 18May2021 (Lot number: ET8885) as dose 1, single for covid-19 immunisation. The mother s relevant medical history and concomitant medications were not reported. Unsure if mother has had symptoms associated with COVID-19. Mother is not currently breastfeeding. The following information was reported: MATERNAL EXPOSURE DURING PREGNANCY (death, congenital anomaly, medically significant) with onset 18May2021, outcome "fatal", described as "Maternal exposure during pregnancy, first trimester"; FOETAL GROWTH RESTRICTION (death, congenital anomaly, medically significant), outcome "fatal", described as "scan at 9 weeks showed fetus stopped growing at 6w 3D"; FOETAL DEATH (death, congenital anomaly, medically significant) with onset 02Jun2021, outcome "fatal" described as "Early miscarriage"; FOETAL CARDIAC ARREST (death, congenital anomaly, medically significant), outcome "fatal", described as "Heart stopped at 6weeks 3 days". The mother had a miscarriage. The fetus"s heart stopped at 6weeks 3 days. Missed miscarriage It was unsure if the medicine have an adverse effect on any aspect of the pregnancy. Mother was exposed to the medicine first-trimester (1-12 weeks). Details of scans or investigations: Vaccine at 3 weeks, scan at 9 weeks showed fetus stopped growing at 6w 3D. The patient underwent the following laboratory tests and procedures: scan: fetus stopped growing, notes: scan at 9 weeks showed fetus stopped growing at 6w 3D. The patient date of death was unknown. The reported cause of death was maternal exposure during pregnancy foetal growth restriction, foetal death, foetal cardiac arrest. It was not reported if an autopsy was performed. No follow-up attempts are possible. No further information is expected.; Sender's Comments: Linked Report(s): GB-PFIZER INC-2021906904 Mother case; Reported Cause(s) of Death: Maternal exposure during pregnancy, first trimester; scan at 9 weeks showed fetus stopped growing at 6w 3D; Early miscarriage; Heart stopped at 6weeks 3 days

Fetus had pulmonary hemorrhage:

<u>CASE 11 (VAERS 2230334)</u> – Pregnant woman had 1st Moderna dose Jul.30, 2021 and 2nd Moderna dose on Aug. 27, 2021. Next day, fetus had pulmonary hemorrhage and peritoneal hemorrhage resulting in stillbirth on Aug. 28, 2021 at 38 weeks gestation.

VAERS ID: 2230334 Vaccinated: 2021-07-30 VAERS Form: 2 Onset: 0000-00-00 Age: Submitted: 0000-00-00 Sex: Unknown Entered: 2022-04-13 Location: Foreign Vaccination / Manufacturer (1 vaccine) Lot / Dose Site / Route COVID19: COVID19 (COVID19 (MODERNA)) / MODERNA 3004672 / 1 Administered by: Unknown Purchased by: ?? Symptoms: Haemoperitoneum, Lymphadenitis, Pulmonary haemorrhage, Foetal death Life Threatening? No Birth Defect? No Died? Yes Date died:2021-08-28 Permanent Disability? No Recovered? No Office Visit (V2.0)? No ER or Office Visit (V1.0)? No ER or ED Visit (V2.0)? No Hospitalized? No **Previous Vaccinations:** Other Medications: Current Illness: Preexisting Conditions: Allergies: Diagnostic Lab Data: CDC 'Split Type': ESMODERNATX, INC.MOD20225 Write-up: This case was received via Regulatory Agency (Reference number: ES-AEMPS-1146166) on 08-Apr-2022 and was forwarded to Moderna on 08-Apr-2022. This regulatory authority case was reported by a consumer (subsequently medically confirmed) and describes the occurrence of PULMONARY HAEMORRHAGE (Haemorrhage pulmonary), HAEMOPERITONEUM (Hemorrhage peritoneal), FOETAL DEATH (FETAL DEATH) and LYMPHADENITIS (Nonspecific mesenteric lymphadenitis) in a neonate of an unknown age and gender exposed to mRNA-1273 (Spikevax) (batch nos. 3005703 and 3004672), while the mother received the product for COVID-19 vaccination. No Medical History information was reported. On 30-Jul-2021, the mother received first dose of mRNA-1273 (Spikevax) (unknown route) 1 dosage form. On 27-Aug-2021, received second dose of mRNA-1273 (Spikevax) (unknown route) dosage was changed to 1 dosage form. Last menstrual period and estimated date of delivery were not provided. On an unknown date, the neonate was diagnosed with PULMONARY HAEMORRHAGE (Haemorrhage pulmonary) (seriousness criterion death), HAEMOPERITONEUM (Hemorrhage peritoneal) (seriousness criterion death), FOÈTAL DEATH (FETAL DEATH) (seriousness criterion death) and LYMPHADENITIS (Nonspecific mesenteric lymphadenitis) (seriousness criterion death). The delivery occurred on an unknown date, which was reported as Unknown. For neonate 1, The outcome was reported as Stillbirth NOS. The neonate died on 28-Aug-2021 An autopsy was performed. The autopsydetermined cause of death was Nonspecific mesenteric lymphadenitis and Hemorrhage pulmonary. mRNA-1273 (Spikevax) (Transplacental)

was withdrawn on 30-Jul-2021. No concomitant and treatment medications were reported. Company Comment: This regulatory case concerns a neonate (age not provided), of unknown gender, with no medical history reported, who experienced the unexpected, serious (fatal) events of foetal death, pulmonary haemorrhage, hemoperitoneum and lymphadenitis. The patient's mother received the first dose of the Moderna mRNA-1273 vaccine on 30Jul2021 and the second dose of the Moderna mRNA-1273 vaccine on 27Aug2021. The patient was exposed to the vaccine at 38 weeks of gestation. The patient expired on 28Aug2021 (1 day after the mother received the second dose). An autopsy was performed and the autopsy-determined causes of death were "Nonspecific mesenteric lymphadenitis" and "Pulmonary hemorrhage". No further details were provided regarding risk factors including infections or drug intake during pregnancy. The benefit-risk relationship of the Moderna mRNA-1273 vaccine is not affected by this report. Sender's Comments: This regulatory case concerns a neonate (age not provided), of unknown gender, with no medical history reported, who experienced the unexpected, serious (fatal) events of foetal death, pulmonary haemorrhage, hemoperitoneum and lymphadenitis. The patient's mother received the first dose of the Moderna mRNA-1273 vaccine on 30Jul2021 and the second dose of the Moderna mRNA-1273 vaccine on 27Aug2021. The patient was exposed to the vaccine at 38 weeks of gestation. The patient expired on 28Aug2021 (1 day after the mother received the second dose). An autopsy was performed and the autopsy-determined causes of death were "Nonspecific mesenteric lymphadenitis" and "Pulmonary hemorrhage" No further details were provided regarding risk factors including infections or drug intake during pregnancy. The benefit-risk relationship of the Moderna mRNA-1273 vaccine is not affected by this report. Autopsy-determined Cause(s) of Death: Nonspecific mesenteric lymphadenitis;

Placenta problems

Hemorrhage pulmonary.

<u>CASE 12 (VAERS 2386294)</u> – 34 year old Pregnant woman had 3rd Pfizer jab on Jan. 20, 2022. Then 20 days later fetus had pulmonary hemorrhage resulting in stillbirth at 34+3 weeks. Cause of death: PLACENTAL FAILURE.

VAERS ID: 2386294 Vaccinated: 2022-01-20 VAERS Form: 2 Onset: 2022-01-20 Submitted: 0000-00-00 Age: Unknown Entered: 2022-07-23 Sex: Location: Foreign Vaccination / Manufacturer (1 vaccine) Lot / Dose Site / Route COVID19: COVID19 (COVID19 (PFIZER-BIONTECH)) / PFIZER/BIONTECH FN5519 3 Administered by: Other Purchased by: ?? Symptoms: Heart disease congenital, Placental insufficiency, Pulmonary congestion, Pulmonary haemorrhage Foetal death, Investigation, Maternal exposure timing unspecified, Specialist consultation, Growth disorder Life Threatening? No Birth Defect? No Died? Yes Date died:0000-00-00 Permanent Disability? No Recovered? No Office Visit (V2.0)? Yes ER or Office Visit (V1.0)? No ER or ED Visit (V2.0)? No Hospitalized? Yes, days: (blank) Extended hospital stay? No Previous Vaccinations: Other Medications: SEROQUEL; SOMAC Current Illness: **Preexisting Conditions:** Allergies: Diagnostic Lab Data: Test Name: special histological examination; Result Unstructured Data: Test Result: Additional information after special; Comments; histological examination of the placenta provides evidence for indicating placental failure as the cause of fetal death. The baby was normal-sized, in fact slightly above average, which does not fit so well with placental abruption. With regard to proven maturation disorders in the placenta; Test Name: Pathologist consultation; Result Unstructured Data: Test Result: Pathologist concludes that the combination; Comments: of small circulatory disturbance placenta and immaturity with poorly developed vascular membranes may explain placental failure.; Test Name: routine growth control; Result Unstructured Data: Test Result results in notes; Comments Uncomplicated pregnancy until she came to routine growth control 09Dec2022 due to her treatment with quetiapine (Seroquel).; Test Date: 20220209; Test Name: routine growth control; Result Unstructured Data: Test Result: results in notes; Comments: Intrauterine fetal death was then detected, gestation week 34+3. No sign of life since the night before. CDC 'Split Type': NOPFIZER INC202200973069
Write-up: HISTOLOGICAL EXAMINATION OF THE PLACENTA PROVIDES EVIDENCE FOR INDICATING PLACENTAL FAILURE AS THE CAUSE OF FETAL DEATH; Congestion and bleeding in the lungs; Congestion and bleeding in the lungs; Dilated right side of heart; Signs of cause of FeTal DEATH, Congestion and bleeding in the range, Congestion and bleeding in the range, Dilated right side of recal, Signs maturation disorders; DEATH INTRAUTERINE; Maternal Drug Exposure; This is a spontaneous report received from a contactable reporter(s) (Physician) from the Regulatory Authority-WFB. A fetus patient was exposed to BNT162b2 (COMIRNATY), transplacental, administration details for the mother: or 20Jan2022 at 12:00 as dose 3 (booster) single (Lot number: FN5519) intramuscular for covid-19 immunisation. The mother of the patient was 34 years old. The mother was 34 weeks pregnant at the event onset Concomitant medication(s) included: SEROQUEL transplacental, start date: 01Jan2021, SOMAC transplacental, start date: 01Jan2022. The following information was reported: MATERNAL EXPOSURE TIMING UNSPECIFIED (death) with onset 20 Jan 2022, outcome "fatal", described as "Maternal Drug Exposure"; FOETAL DEATH (medically significant) with onset 09Feb2022, outcome "unknown", described as "DEATH INTRAUTERINE"; PULMONARY HAEMORRHAGE (death, medically significant), PULMONARY CONGESTION (death, medically significant), outcome "fatal" and all described as "Congestion and bleeding in the lungs"; HEART DISEASE CONGENITAL (death, medically significant), outcome "fatal", described as "Dilated right side of heart"; GROWTH DISORDER (death), outcome "fatal", described as "Signs of maturation disorders": PLACENTAL INSUFFICIENCY (hospitalization, medically significant), outcome "unknown", described as "HISTOLOGICAL EXAMINATION OF THE PLACENTA PROVIDES EVIDENCE FOR INDICATING PLACENTAL FAILURE AS THE CAUSE OF FETAL DEATH". The event "death intrauterine" required physician office visit. The patient underwent the following laboratory tests and procedures: special histological examination: (unspecified date) Additional information after special, notes: histological examination of the placenta provides evidence for indicating placental failure as the cause of fetal death. The baby was normal-sized, in fact slightly above average, which does not fit so well with placental abruption. With regard to proven maturation disorders in the placenta; Specialist consultation. (unspecified date) Pathologist concludes that the combination, notes: of small circulatory disturbance placenta and immaturity with poorly developed vascular membranes may explain placental failure; (unspecified date) results in notes, notes. Uncomplicated pregnancy until she came to routine growth control 09Dec2022 due to her treatment with quetiapine (Seroquel); (09Feb2022) results in notes, notes Intrauterine fetal death was then detected, gestation week 34+3. No sign of life since the night before. The patient date of death was unknown. Reported cause of death: "Congestion and bleeding in the lungs". Clinical course: Finds no pathological changes in the fetus other than congestion and bleeding in the lungs. Dilated right side of heart, but normal heart growth and no convincing signs of cardiomyopathy or myocarditis. Bleeding perceived as asphyxia conditioned. The placenta is described as small and with maternal circulatory disturbance. Simultaneous signs of maturation disorders that are sometimes seen in diabetes during pregnancy. Pathologist concludes that the combination of small circulatory disturbance placenta and immaturity with poorly developed vascular membranes may explain placental failure. Additional information after special histological examination of the placenta provides evidence for indicating placental failure as the cause of fetal death. The baby was normal-sized, in fact slightly above average, which does not fit so well with placental abruption. With regard to proven maturation disorders in the placenta. Causal relationship between the event Death intrauterine and the administration of CORMINATY was assessed as Possible by Regulatory Authority; Causal relationship between the event Placental insufficiency and the administration of Quetiapine was assessed as Possible by Regulatory Authority. No follow-up attempts are possible. No further information is expected.; Sender's Comments: Linked Report(s): NO-PFIZER INC-202200618967 fetus report;; Reported Cause(s) of Death: Congestion and bleeding in the lungs

<u>CASE 13 (VAERS 2046121)</u> – Pregnant woman had 2nd dose of Moderna on Oct.20, 2021 and 6 days later fetus died due to "clotting at the placenta" and fetal vascular malperfusion. Reported cause of death: clotting at the placenta.

VAERS ID: 2046121 Vaccinated: 2021-09-22 VAERS Form: 2 Onset: 0000-00-00 Submitted: 0000-00-00 Age: Unknown Entered: 2022-01-19 Sex: Location: Vaccination / Manufacturer (1 vaccine) Lot / Dose Site / Route COVID19: COVID19 (COVID19 (MODERNA)) / MODERNA 062E21A 2 Administered by: Unknown Purchased by: ?? Symptoms: Foetal death, Foetal vascular malperfusion Life Threatening? No Birth Defect? No Died? Yes Date died: 2021-10-26 Permanent Disability? No Recovered? No Office Visit (V2.0)? No ER or Office Visit (V1.0)? No ER or ED Visit (V2.0)? No Hospitalized? No Previous Vaccinations: Other Medications: Current Illness: **Preexisting Conditions:** Allergies: Diagnostic Lab Data: CDC 'Split Type': USMODERNATX, INC.MOD20224 Write-up: clotting at the placenta/caused the baby to not receive any blood. Baby was deceased on 26Oct2021. This spontaneous case was reported by a consumer and describes the occurrence of FOETAL VASCULAR MALPERFUSION (clotting at the placenta/caused the baby to not receive any blood) and FOETAL DEATH (Baby was deceased on: 26Oct2021) in a foetus patient of an unknown gender who received mRNA-1273 (Moderna COVID-19 Vaccine) (batch nos. 062E21A and 026D21D) for COVID-19 vaccination. MEDICĀL HISTORY (Parent): The mother"s past medical history included Maternal exposure during pregnancy. No Medical History information was reported. On 22-Sep-2021, the patient received first dose of mRNA-1273 (Moderna COVID-19 Vaccine) (Transplacental) 1 dosage form On 20-Oct-2021. received second dose of mRNA-1273 Moderna COVID-19 Vaccine) (Transplacental) dosage was changed to 1 dosage form. On an unknown date, the patient experienced FOETAL VASCULAR MALPERFUSION (clotting at the placenta/caused the baby to not receive any blood) (seriousness criteria death and medically significant) and FOETAL DEATH (Baby was deceased on: 26Oct2021) (seriousness criteria death and medically significant). The patient died on 26-Oct-2021 The reported cause of death was clotting at the placenta It is unknown if an autopsy was performed. No concomitant and treatment medication were provided Patient stated that this is my "rainbow baby" as her previous baby passed away due to doctor negligence in 2019 as she stated that it had nothing to do with health it was doctor"s negligence in 2019. Company Comment: This case refers to a foetal patient of unspecified age and gender with a medical history of mother reporting maternal exposure during pregnancy. The patient experienced the unexpected event of Foetal vascular malperfusion on an unspecified date after the second dose of mRNA-1273 vaccine but experienced the unexpected event of Foetal death approximately 6 days after receiving the second dose of the vaccine. No causality assessment was provided by the reporter. The benefit-risk relationship of mRNA-1273 is not affected by this report. This case was linked to MOD-2022-440025 (Patient Link); Sender"s Comments: This case refers to a foetal patient of unspecified age and gender with a medical history of mother reporting maternal exposure during pregnancy. The patient experienced the unexpected event of Foetal vascular malperfusion on an unspecified date after the second dose of mRNA-1273 vaccine but experienced the unexpected event of Foetal death approximately 6 days after receiving the second dose of the vaccine. No causality assessment was provided by the reporter. The benefit-risk relationship of mRNA-1273 is not affected by this report.; Reported Cause(s) of Death: Clotting at the placenta

<u>CASE 14 (VAERS 2463841)</u> – 30 year old pregnant woman had 3rd Pfizer jab on Dec. 18, 2021 and one month later on Jan. 21, 2022 ultrasound showed growth restriction, on Feb.20, 2022 fetus had no heartbeat. Stillbirth on Feb. 23, 2022. Cause of death: placental insufficiency caused by severe maternal vascular malperfusion.

VAERS ID: 2463841 Vaccinated: 2021-12-18 VAERS Form: 2 Onset: 2021-12-18 Age: Submitted: 0000-00-00 Sex: Female Entered: 2022-09-29 Location: Vaccination / Manufacturer (1 vaccine) Lot / Dose Site / Route COVID19: COVID19 (COVID19 (PFIZER-BIONTECH)) / PFIZER/BIONTECH Administered by: Other Purchased by: ?? Symptoms: Foetal heart rate abnormal, Foetal growth restriction, Maternal exposure during pregnancy Life Threatening? No Birth Defect? No Died? Yes Date died:2022-02-23 Permanent Disability? No. Recovered? No Office Visit (V2.0)? No ER or Office Visit (V1.0)? No ER or ED Visit (V2.0)? No Hospitalized? No Previous Vaccinations: Other Medications: ELTROXIN; FOLIC ACID Current Illness: Preexisting Conditions: Allergies: Diagnostic Lab Data: CDC 'Split Type': IEPFIZER INC202201191407 Write-up: The baby presented with an absent fetal heartbeat; growth restriction/ fetal growth restriction on a recent ultrasound/ intrauterine growth restriction; Maternal exposure during pregnancy; This is a spontaneous report received from contactable reporter(s) (Consumer or other non HCP and Physician) from the RA. A female patient was exposed to BNT162b2 (COMIRNATY), transplacental, administration details for the mother: on 18Dec2021 as dose 3 (booster) single (Batch/Lot number: unknown) for covid-19 immunisation. The mother of the patient was 30 years old. The patient's relevant medical history was not reported. The mother's relevant medical history included: "Subclinical hypothyroidism", start date: 08Jun2020 (ongoing); "Verruca", start date: 19Oct2021 (unspecified if ongoing), notes: x2, advised re filing/salicylic acid; "SUBDERMAL HYPOTHYROIDISM / SUBCLINICAL HYPOTHYROIDISM" (unspecified if ongoing). The mother was 26 weeks pregnant at the event onset. Concomitant medication(s) included: ELTROXIN transplacental taken for hypothyroidism, start date: 02Feb2020, stop date: 18Aug2022; FOLIC ACID transplacental. The mother s vaccination history included: BNT162b2 (DOSE 1, SINGLE, Lot No.FCS029, Event start date: Jul2021, Outcome: Not Recovered), administration date: 20Jul2021, for Covid-19 immunization reaction(s): "red rash across torso/ At first the rash was oval in shape, reddish with a yellow centre. Over the following months it has changed in shape and is now just red blotches", "discus lesion/ Some new lesions on lower abdo wall / one new lesion is pink, discoid and blanches / discoid rash / right abdominal wall has pale pink discoid superficial dermatitis lesions. These were originally "targetoid" but seem to have settle", "Prickly heat"; comirnaty (dose 2, single (Lot number: FE7053), administration date: 10Aug2022, for Covid-19 immunization, reaction(s): "the patient developed discoid lesions on her abdomen wall after her first COVID-19 vaccine and this then flared after her second vaccine." The following information was reported: MATERNAL EXPOSURE DURING PREGNANCY (death) with onset 18Dec2021. outcome "fatal" FOETAL GROWTH RESTRICTION death) with onset 21Jan2022, outcome "fatal", described as "growth restriction/ fetal growth restriction on a recent ultrasound/intrauterine growth restriction"; FOETAL HEART RATE ABNORMAL (death) with onset 20Feb2022, outcome "fatal", described as "The baby presented with an absent fetal heartbeat". The pregnancy resulted in still birth. The patient date of death was 23Feb2022 Reported cause of death: "The baby presented with an absent fetal heartbeat", "growth restriction/ fetal growth restriction on a recent ultrasound/ intrauterine growth restriction". No autopsy was performed. Clinical course: A placental exam showed the cause of death was placental insufficiency, caused by maternal vascular malperfusion. They did not have an autopsy. On 20Feb2022, the baby presented with an absent fatal heartbeat related to this severe placental disease. A placental examination showed a hyper coiled cord but more importantly, severe maternal vascular malperfusion. On 21Jan2022, it was noted there was fatal growth restriction on a recent ultrasound. No follow-up attempts are possible; information about lot/batch number cannot be obtained. No further information is expected.; Sender's Comments: Linked Report(s): IE-PFIZER INC-202201137917 Same Product, Different Dose/Patient/Event (Dose 1);IE-PFIZER INC-202201136145 Same Product, Different Dose/Patient/Event (Dose 3);IE-PFIZER INC-202201191408 Same Product, Different Dose/Patient/Event (Dose 2); Reported Cause(s) of Death: The baby presented with an absent fetal heartbeat; growth restriction/ fetal growth restriction on à recent ultrasound/ intrauterine growth restriction

My Take...

Every pregnant woman should have had these risks explained to her as part of informed consent. The risks of Pfizer and Moderna COVID-19 mRNA vaccines to the fetus include:

- fetus can stop growing and die immediately after taking the mRNA vaccine or up to several days, weeks or months later, resulting in miscarriage or stillbirth;
- fetus can suffer cardiac arrest and die;
- fetus can suffer pulmonary hemorrhage and die;
- fetus can die from placental blood clots
- fetus can die from placental insufficiency or placental failure
- fetus can die from congenital malformations (next substack)

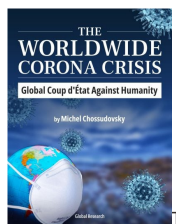
Please note that these COVID-19 mRNA vaccine related adverse events were known and recorded in VAERS in early 2021 so every doctor should have been advising their pregnant patients about these documented and known adverse events.

Unfortunately, the vast majority of doctors did not educate their pregnant patients about the risks of Pfizer and Moderna COVID-19 mRNA vaccines, and should be held legally liable for failing to do so.

*

Note to readers: Please click the share button above. Follow us on Instagram and Twitter and subscribe to our Telegram Channel. Feel free to repost and share widely Global Research articles.

Dr. William Makis is a Canadian physician with expertise in Radiology, Oncology and Immunology. Governor General's Medal, University of Toronto Scholar. Author of 100+ peer-reviewed medical publications.



The Worldwide Corona Crisis, Global Coup d'Etat Against

Humanity

by Michel Chossudovsky

Michel Chossudovsky reviews in detail how this insidious project "destroys people's lives". He provides a comprehensive analysis of everything you need to know about the "pandemic" — from the medical dimensions to the economic and social repercussions, political underpinnings, and mental and psychological impacts.

"My objective as an author is to inform people worldwide and refute the official narrative which has been used as a justification to destabilize the economic and social fabric of entire countries, followed by the imposition of the "deadly" COVID-19 "vaccine". This crisis affects humanity in its entirety: almost 8 billion people. We stand in solidarity with our fellow human beings and our children worldwide. Truth is a powerful instrument."

ISBN: 978-0-9879389-3-0, Year: 2022, PDF Ebook, Pages: 164, 15 Chapters

We encourage you to support the eBook project by making a donation through Global Research's <u>DonorBox "Worldwide Corona Crisis" Campaign Page</u>.

The original source of this article is <u>COVID Intel</u> Copyright © <u>Dr. William Makis</u>, <u>COVID Intel</u>, 2023

Comment on Global Research Articles on our Facebook page

Become a Member of Global Research

Articles by: Dr. William Makis

Disclaimer: The contents of this article are of sole responsibility of the author(s). The Centre for Research on Globalization will not be responsible for any inaccurate or incorrect statement in this article. The Centre of Research on Globalization grants permission to cross-post Global Research articles on community internet sites as long the source and copyright are acknowledged together with a hyperlink to the original Global Research article. For publication of Global Research articles in print or other forms including commercial internet sites, contact: publications@globalresearch.ca

www.globalresearch.ca contains copyrighted material the use of which has not always been specifically authorized by the copyright owner. We are making such material available to our readers under the provisions of "fair use" in an effort to advance a better understanding of political, economic and social issues. The material on this site is distributed without profit to those who have expressed a prior interest in receiving it for research and educational purposes. If you wish to use copyrighted material for purposes other than "fair use" you must request permission from the copyright owner.

For media inquiries: $\underline{publications@globalresearch.ca}$