Histopathologically TMA-like distribution of multiple organ thromboses following the initial dose of the BNT162b2 mRNA vaccine (Comirnaty, Pfizer/BioNTech): an autopsy case report

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Abstract

Background  Coronavirus disease 2019 (COVID-19) has spread worldwide. Vaccination is now recommended as one of the effective countermeasures to control the pandemic or prevent the worsening of symptoms. However, its adverse effects have been attracting attention. Here, we report an autopsy case of multiple thromboses after receiving the first dose of the BNT162b2 mRNA vaccine (Comirnaty, Pfizer/BioNTech) in an elderly woman.

Case presentation  A 72-year-old woman with a history of diffuse large B-cell lymphoma in the stomach and hyperthyroidism received the first dose of the BNT162b2 mRNA vaccine and died 2 days later. The autopsy revealed multiple microthrombi in the heart, brain, liver, kidneys, and adrenal glands. The thrombi were CD61 and CD42b positive and were located in the blood vessels primarily in the pericardial aspect of the myocardium and subcapsular region of the adrenal glands; their diameters were approximately 5–40 μm. Macroscopically, a characteristic myocardial haemorrhage was observed, and the histopathology of the characteristic thrombus distribution, which differed from that of haemolytic uraemic syndrome and disseminated intravascular coagulation, suggested that the underlying pathophysiology may have been similar to that of thrombotic microangiopathy (TMA).

Conclusion  This is the first report on a post-mortem case of multiple thromboses after the BNT162b2 mRNA vaccine. The component thrombus and characteristic distribution of the thrombi were similar to those of TMA, which differs completely from haemolytic uraemic syndrome or disseminated intravascular coagulation, after vaccination. Although rare, it is important to consider that fatal adverse reactions may occur after vaccination and that it is vital to conduct careful follow-up.

Keywords  SARS-CoV-2, Coronavirus disease 2019, Vaccination, BNT162b2, Thrombotic microangiopathy, Thrombosis

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Background
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is a novel virus that was first reported in late 2019 as the causative agent of coronavirus disease 2019 (COVID-19), which has spread worldwide [1]. The risk of developing severe COVID-19 symptoms is particularly high among the elderly and individuals with underlying diseases [2]. One of the effective countermeasures to control the pandemic or prevent the worsening of symptoms, as dictated by the World Health Organization, is vaccination. With the rapid implementation of vaccination programmes and a subsequent increase in the total number of vaccinated individuals, adverse vaccine effects have been attracting attention. Serious side effects such as (peri) myocarditis following mRNA-based vaccination [3–5] and thrombosis following adenoviral vector-based vaccination [6–8] have been reported. Although a number of post-mortem investigations of fatalities following mRNA vaccination showed myocardial infarction or myocarditis [9], autopsy reports showing thrombotic microangiopathy (TMA)-like multiple organ thromboses following mRNA vaccination are rare. Here, we report an autopsy case of a patient who died from systemic thrombosis within 2 days of receiving the BNT162b2 mRNA vaccine (Comirnaty, Pfizer/BioNTech).

Case presentation
A 72-year-old woman with a history of hyperthyroidism, an unspecified penicillium allergy developed in her 20s, and diffuse large B-cell lymphoma in the stomach received the first dose of the BNT162b2 mRNA vaccine at about 9 a.m. on day X. Her pre-vaccination screening questionnaire for the COVID-19 vaccine showed that her body temperature on the vaccination day was 35°C, and no other notable findings were described. Her lymphoma showed a complete response to chemotherapy with an unexpected course, and her hyperthyroidism was under control with oral therapy. She had no medical history of deep vein thrombosis, systemic lupus erythematosus, recurrent pregnancy loss, haematuria, and haematopoietic stem cell or solid organ transplantation. Laboratory testing showed that her liver and kidney function, as well as blood count, were within normal limits one month before vaccination. The absence of thrombocytopenia and anaemia was confirmed 10 days before vaccination. She felt unwell at the vaccination venue immediately after the vaccination; however, her condition improved with some rest after which she went home. About 4 p.m. on day X+1, she developed fatigue, nausea, chest pain, and back pain, and around 8 p.m. on day X+1 LINE, a social networking service, showed a “Read” mark suggesting that she was alive. She was found deceased in her house on day X+2. Life-saving procedures were not administered because rigor mortis was noted in the muscles of the jaw. A medicolegal autopsy was performed approximately 24 h after she was found deceased to investigate the manner of her death, since vaccination was suspected to have been the cause.

Autopsy findings
At autopsy, the patient’s body length and weight were 155 cm and 53.0 kg, respectively. There were no reddening or wheals observed on the body surface. The heart weight was 394 g, and 170 mL of concentrated yellow-translucent pericardial fluid with fibrinous precipitate was present in the pericardium. Marked petechial haemorrhage was found on the surface of the posterior pericardium (Fig. 1a). Gross examination after 10% buffered formalin fixation revealed black-red discoloration throughout the circumference of the pericardium and outer surface of the myocardium (Fig. 1b). The left anterior descending artery showed only 25% angiostenosis, and no evident obstruction was noted in the coronary arteries. Marked petechial haemorrhage was also noted on the surface of the lungs, liver, kidneys, and spleen (Additional file 1a); on the diaphragm, lateral and posterior pleura; and on the mucosal surfaces of the oesophagus (Additional file 1b), stomach, and duodenum. An additional picture file shows haemorrhagic lesion in the spleen and oesophagus [see Additional file 1]. Haemorrhagic lesions were not observed in the intracranial space, and thrombosis was not noted in the superior sagittal sinus. The bladder urine was not bloody. There were no findings associated with traumatic injuries.

Histopathological findings
Histopathological examination revealed multifocal vacuolation and lipofuscin pigmentaton in cardiomyocytes. Immunohistochemical (IHC) staining showed positivity for anti-C4d antibodies (rabbit polyclonal, catalogue no. 12-5000; American Research Products, Inc., Belmont, MA, USA; 1:400) in the vacuolated cells of the left ventricular free wall (Additional file 2a and 2b). An additional picture file shows anti-C4d -positive vacuolated cells [see Additional file 2]. Numerous microthrombi without inflammatory cells were found in the small vessels, arterioles, and capillaries of the anterior, posterior, and lateral walls of the left ventricle, right ventricle, and interventricular septum, located predominantly at the border between the haemorrhagic and non-haemorrhagic areas (Fig. 1c and f). These microthrombi were found to be immunoreactive for anti-CD42b (mouse monoclonal, clone MM2/174; Novocastra, Newcastle upon Tyne, UK; 1:100), anti-CD61 (mouse monoclonal, clone 2F2; Leica Biosystems Newcastle, Ltd., Newcastle upon Tyne, UK; 1:500), and anti-von Willebrand factor (vWF) (mouse monoclonal, clone F8/86; Dako, Carpinteria, CA, USA; 1:25) antibodies, as well as periodic acid

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Fig. 1 Macroscopic and microscopic findings of the heart. 

(a) Macroscopic haemorrhage in the posterior pericardium in situ at the autopsy. 

(b) Gross findings of the heart after fixation. The cut surface of the heart after fixation shows black-red discoloration (arrowhead) in the entire circumference of the pericardium and pericardium-side myocardium.

(c) Scanning magnifications of the heart with haematoxylin-eosin (HE) staining.

(d) Schematic illustration of the microscopic pathology. The red dots indicate the microscopic haemorrhage, the blue rectangle indicates the thrombus, and the green arc indicates the contraction band necrosis. 

(LV; left ventricle, IVS; interventricular septum, RV; right ventricle.

(e) and (f) Low- and high-power views of the haemorrhage in the cardiomyocytes. The haemorrhages were found in the pericardium and pericardium-side myocardium, which is compatible with the discoloration in macroscopic observation. Scale bars indicate 100 μm (e, f)
Schiff staining (Fig. 2a and e). An additional picture file shows ultrastructural analysis of a thrombus in the heart in detail [see Additional file 3]. The vascular diameter of the microthrombi was approximately 5–40 μm, and the microvessels were congested and dilated. Increased eosinophilic cardio-myofibrillar bundles were localised,
and these bundles were highlighted with Luxol Fast Blue (LFB) stain. LFB staining also demonstrated many irregular deep-blue wavy contraction bands extending across the fibres in the region of the cardiomyocytes, indicating contraction band necrosis. No significant inflammatory cell infiltration or fibrotic lesion was detected in the cardiac specimen. In addition, IHC for anti-SARS-CoV-2 spike glycoprotein (rabbit monoclonal, clone HL6; GeneTex, Inc., Irvine, CA, USA; 1:100) was negative in the cardiomyocytes.

The kidneys were found to have focal interstitial haemorrhage, as well as fibrin and platelet thrombi in the glomerular capillaries and afferent/afferent arterioles. CD61-positive thrombi were detected in the small vessels and arteriole of the renal subcapsular region. Although IHC staining showed positivity for anti-C4c antibodies partially in the glomerular tuft and capillaries, as well as the parenchyma of the renal tubule, it was difficult to assess. Jones sliver (periodic acid-methenamine silver) staining revealed the enlarged subendothelial space of the glomerular basement membrane in some regions of the glomerulus. Microthrombi were also found in the liver, pancreas, cerebral cortex, cerebellum, and pons; however, no ischaemic change was noted. Thrombi covered with endothelium were not detected in any of the tested organs. The vascular diameter of the vessels involved in thrombi formation in these organs was similar to that of those in the heart. In the lungs, the pulmonary capillaries were congested, and pulmonary oedema was noted. Haemorrhagic lesions were found in the capsule and parenchyma of the spleen and lateral lacunae of the superior sagittal sinus. There were no apparent findings associated with infections such as inflammatory cell infiltration or autoimmune diseases including systemic sclerosis and nephropathy.

Virus isolation tests did not detect any viruses in the pericardial fluid. Her blood was not biochemically tested for factors such as autoantibodies, a disintegrin-like and metalloprotease with thrombospondin type 1 motifs, member 13 (ADAMTS13) activities, and platelets, because of post-mortem changes at the time of autopsy.

As mentioned above, platelet microthrombi were detected in multiple organs, predominantly in the heart, and injuries to the other organs were limited. Therefore, it was conceivable that the sudden death was cardiac in origin.

**Discussion and conclusions**

Since March 2021, unusual thrombosis and thrombocytopenia have been reported after receiving the ChAdOx1 nCoV-19 vaccine, an adenoviral vector-based vaccine (Oxford/AstraZeneca) [6–8]. Novel pathological concepts, namely vaccine-induced immune thrombotic thrombocytopenia (VITT) [10], vaccine-induced prothrombotic immune thrombocytopenia (VIPIT) [11], and thrombosis with thrombocytopenia syndrome (TTS) [12, 13] were proposed as the underlying mechanisms. Research on thrombosis post-vaccination has revealed an association between VITT and the heparin-induced thrombocytopenia (HIT) antibody [7, 14]. Thrombotic events can also be caused by mRNA vaccines, although only a few reports are available and the event frequency after mRNA-based vaccination is lower than after adenovirus vector vaccination [15–18]. In our case, the deceased was found dead 2 days after receiving the mRNA vaccine and her death was suggestive of a vaccine-associated death.

In our case, there were four characteristic macroscopic/microscopic findings: (i) thrombi distributed to most of the systemic organs, especially the heart, (ii) thrombi located primarily in the pericardium-side myocardium and subcapsular regions of the adrenal gland, (iii) thrombi formations 5–40 μm in size found in capillaries and arterioles, and (iv) morphology and IHC staining revealed that the thrombi primarily comprised platelets. Platelet thrombi in the glomerular capillaries and the enlarged subendothelial space of the glomerular basement membrane suggested a vascular endothelial injury in the acute phase. Platelet-rich thrombi are formed by excessive activation of platelet or the vascular endothelial injury and, in our case, the vascular endothelial injury was mainly involved. The absence of thrombi covered with endothelium could be an indication that the thrombi were also in the acute phase. Although laboratory data in the post-mortem investigation were absent, these histopathological findings were compatible with TMA, especially TTP [19–23]; thus, we hypothesised that the underlying pathophysiology in our case was similar to TMA. In this context, TMA meant “histopathological TMA” because TMA is a clinicopathological concept including the triad microangiopathic haemolytic anaemia, thrombocytopenia, and organ dysfunction due to thrombi formation in small vessels. The differential diagnosis of systemic thrombosis involves disseminated intravascular coagulation (DIC), paroxysmal nocturnal haematuria (PNH), autoimmune HIT, TTS/VITT, and TMA. The concept of TMA includes a broad spectrum of diseases [24], such as thrombotic thrombocytopenic purpura (TTP) occurring after adenoviral vector-based vaccination and haemolytic uraemic syndrome (HUS), and TMA with autoimmune conditions, for instance, catastrophic antiphospholipid syndrome (CAPS). Although it is impossible to make a complete differentiation based on each definition and criteria of each disease due to the lack of clinical symptoms, detailed history, and laboratory data including autoantibodies, complement titres, and blood counts, we attempted to histopathologically compare the differentiation of each disease based on
the macro/microscopic findings obtained from autopsies and/or the clinically confirmed thrombotic lesion (Table 1). In HUS, thrombi are found primarily in the kidneys [19]. DIC characteristically presents with primarily fibrin thrombi in the renal glomerular capillaries [25, 26], whereas our case presented with platelet thrombi in the afferent/efferent arteriole-level vessels of multiple organs. According to laboratory data including LDH levels one month before the vaccination, the blood count 10 days before the vaccination, and no history or autopsy findings of haematuria, our case seemed incompatible with PNH. Moreover, thrombosis at the hepatic vein is characteristic in PNH [27] and haemosiderin deposition is observed in the renal tubular endothelium in chronic cases [28, 29].

Among the few autopsy case reports that include histopathological analysis of HIT [30–33], the lung and gastrointestinal tract are reported as the main thrombosis sites in HIT. TTS/VITT, which is suspected to be associated with autoimmune HIT, develops 4 to 28 days after vaccination and causes thrombosis characteristically in atypical sites such as the cerebral venous sinus, hepatic vein, and splenic vein [34–40]. Although the thrombi in CAPS share an organ distribution similar to that of our case [41–44], thrombi in CAPS form regardless of the shear stress in arterioles and/or capillaries; thus, it might be difficult to explain our four characteristics findings. As mentioned above, the histopathological findings in our case were different from those of these diseases, although we could not analyse the blood; moreover, the history of trauma and infection that could cause autoimmune HIT are also unknown.

TMA-like pathophysiology possibly caused severe consumption thrombocytopenia, resulting in multiple petechial haemorrhages in many organs, such as the pancreas and oesophagus, as observed in our case. However, since blood tests were not available at the time of autopsy, it is difficult to determine whether the vaccination caused or triggered her condition resulting in death. In contrast to a number of post-mortem investigations of fatalities following mRNA vaccination that showed myocardial infarction or myocarditis, our histopathological investigation indicated that multiple fatal thromboses following mRNA-based vaccination may have occurred. Macroscopic haemorrhage was also noted on autopsy, consistent with the region in which the deceased complained of pain. Clinicians and researchers should bear in mind that thrombosis may also be one post-mRNA-based vaccination reaction.

| Pathophysiology          | Characteristic thrombi distribution                                                                 | Chief thrombus vessel                  | Thrombus components | Other features (if any)                                    | Reference |
|--------------------------|--------------------------------------------------------------------------------------------------------|----------------------------------------|---------------------|----------------------------------------------------------|-----------|
| TTP                      | Heart, adrenal gland, kidney, pancreas, brain, liver, and spleen                                      | Arteriole/capillary                   | Platelet            | Thrombi were rarely found in the lung, bone marrow, and GI tract | [19–22]  |
| HUS                      | Kidney and pancreas                                                                                   | Arteriole/capillary                   | Fibrin              |                                                          | [19]     |
| TMA with catastrophic APS| Kidney, heart, lung, brain, and spleen                                                                 | Arteriole/capillary                   | Platelet/Fibrin     | Characteristics of vascular endothelial injuries and proliferation on small arterioles | [41–44]  |
| TMA with SSc             | Kidney (intracapillary)                                                                             | Arteriole/capillary                   | Fibrin              |                                                          | [49, 50] |
| TMA after bone marrow transplant | Kidney and intestine                                                                            | Arteriole/small arteries              | Fibrin              | In the large intestine, the right colon was affected      | [51–53]  |
| Pregnancy associated TMA, HELLP | Kidney, lung, and intestine                                                                              | Arteriole                             | Platelet/Fibrin     | Multiple blackish-reddish patches and confluent hemorrhagic foci on the liver | [54–56]  |
| DIC                      | Kidney, lung, spleen, and adrenal gland                                                               | Arteriole/capillary                   | Fibrin              |                                                          | [25, 26] |
| PNH                      | Hepatic vein                                                                                        | Veins > arteries                      | Fibrin/platelet     | Hemosiderin deposition on the renal tubular epithelium    | [27–29]  |
| HIT                      | Lung, GI tract                                                                                       | Arteries/veins                        | Platelet            |                                                          | [30–33]  |
| TTS/VITT                 | CVS, portal, splenic, and SMV                                                                         | Veins                                 | Platelet/Fibrin     |                                                          | [34–40]  |
| TMA after mRNA-1273 vaccination | Kidney, liver, and GI tract                                                                            | Small arteries                        | Platelet            | Lupus anticoagulant was positive                          | [46]     |
| Our case                 | Heart, adrenal gland, kidney, liver, pancreas, brain, and pons                                        | Arteriole/capillary                   | Platelet            | Thrombi were not found in the lungs or GI tract           | [47]     |

TMA=trombotic microangiopathy, TTP=trombotic thrombocytopenic purpura, HUS=haemolytic uraemic syndrome, APS=antiphospholipid syndrome, SSc=systemic sclerosis, HELLP=haemolysis, elevated liver function tests, low platelets syndrome, DIC=disseminated intravascular coagulation, PNH=paroximal nocturnal haemoglobinuria, HIT=heparin-induced thrombocytopenia, TTS=trombosis with thrombocytopenia syndrome, VITT=vaccine-induced immune thrombotic thrombocytopenia, CVS=cerebral venous sinus, GI=gastrointestinal, SMV=superior mesenteric vein.
Several potential mechanisms that cause vaccine-induced adverse effects have been proposed. For example, TTS/VITT after adenoviral vector-based vaccination is considered to have the same pathophysiology via the anti-platelet factor 4 antibodies [14]. On the other hand, the potential mechanisms of thrombosis after mRNA vaccination remain unclear. One possible mechanism proposed by Trougakos et al. is the spike hypothesis, which states that spike glycoprotein induced by vaccination acts as if it were the spike glycoprotein produced during SARS-CoV-2 infection [45]. In an autopsy case report of TMA after mRNA-1237 vaccination, the author indicated the histopathological similarities of the autopsied case to microvascular injuries of COVID-19 via a complement pathway [46]. On the other hand, cardiac histopathology in lethal cases of COVID-19 reveals haemorrhage in the pericardium, congested microvascu- lature with/without microhaemorrhage, and hyaline thrombi [47], which are also consistent with our case findings. Considering that the same histopathological changes were observed after viral infection and vaccination, these changes might be related to the common denominator between viral infection and vaccination, namely, the SARS-CoV-2 spike protein and antibodies elicited against it.

TMA-like systemic thrombosis after vaccination itself is a very rare reaction and each case report only indicated temporal relationships; thus, the cause-effect relationship is still unclear. The risk of COVID-19 worsening is thought to be higher than the risk of adverse effects after vaccination. In contrast, a report comparing thrombotic events after influenza vaccination to those after SARS-CoV-2 vaccination showed a significant increase in thrombotic events after COVID-19 vaccination [48]. More rigorous follow-up after vaccination is needed and it is important to detect symptomatic cases early on to allow for early treatment.

In conclusion, this is a rare report of a post-mortem case showing TMA-like multiple systemic thrombi post BNT162b2 mRNA vaccination with a detailed histopathological analysis. Macroscopically, characteristic myocardial haemorrhage was observed, and the histopathology of the characteristic thrombi distribution was similar to that of TMA. Although rare, it is important to keep in mind that fatal adverse reactions may occur after vaccination, and careful follow-up is important after vaccination.

**List of abbreviations**

| Abbreviation | Full Form |
|--------------|-----------|
| SARS-CoV-2   | severe acute respiratory syndrome coronavirus 2. |
| COVID-19     | coronavirus disease 2019. |
| mRNA         | messenger ribonucleic acid. |
| TMA          | thrombotic microangiopathy |
| IHC          | immunohistochemical. |
| VWF          | von Willebrand factor. |
| LFB          | luxol fast blue |
| ADAMTS13     | a disintegrin-like and metalloproteinase with thrombospondin type 1 motifs, member 13. |
| VITT         | vaccine-induced immune thrombotic thrombocytopenia. |
| VPRIT        | vaccine-induced prothrombotic immune thrombocytopenia. |
| TTS          | thrombosis with thrombocytopenia syndrome. |
| HIT          | heparin-induced thrombocytopenia. |
| TTP          | thrombotic thrombocytopenic purpura. |
| HUS          | haemolytic uraemic syndrome. |
| DIC          | disseminated intravascular coagulation. |
| PNH          | paroxysmal nocturnal haematuria. |
| CAPS         | catastrophic antiphospholipid syndrome. |

**Supplementary Information**

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**Supplementary Material 1:** Macroscopic haemorrhage of the spleen and oesophagus.tif. a: macroscopic haemorrhage on the surface of the spleen. b: macroscopic haemorrhage on the mucosal surface of the oesophagus

**Supplementary Material 2:** Vacuolation and anti-C4d-positive cardiomyocytes.tif. a: Vacuolation of the cardiomyocytes. b: anti-C4d immunohistochemistry-positive cardiomyocytes. The positivity is compatible with the vacuolation

**Supplementary Material 3:** Ultrastructural analysis of microthrombi in the heart.tif. a to d: Microthrombi scanned by transmission electron microscopy. The nuclei of vascular endothelial cells and red blood cells are visible, and platelets and fibrin are found as the boundary indistinct area around the red blood cells with low electron density. The degree of occlusion varies; however, almost all thrombi are non-occlusive. The yellow arrowheads indicate the nuclei of the endothelium, the arrows indicate platelets, the stars indicate fibrin, and the red arrowheads indicate the erythrocytes

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**Authors’ contributions**

RK, KK, KM, and SM performed the medicolegal autopsy; RK, HN, MT, TY, and SY prepared histopathological specimen sections; RK, HN, TU, KH, YI, and KA made the pathological diagnosis; RK wrote the manuscript; and AN, TD, and SM revised and edited the manuscript. All authors read and approved the final manuscript.

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**Data availability**

The datasets obtained and analysed in the current study are available from the corresponding author upon reasonable request.

**Declarations**

**Ethics approval and consent to participate**

Our study adhered to the Ethical Guidelines for Medical and Health Research Involving Human Subjects established by the Government of Japan. Although case reports do not require ethics approval in Japan, ethics approval was obtained from Oita University’s Ethics Committee under the reference number 2282. Informed consent to participate was obtained from the patient’s relatives.

**Consent for publication**

Written informed consent was obtained from the patient’s relatives to publish this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.
Competing interests
The authors declare that they have no competing interests.

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