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ABSTRACT

The effective value of microbiological post-mortem examinations stands as fundamental in forensic cases involving microbiology. We ran these analyses on five victims, who suddenly died after showing persistent fever. The examinations were conducted between 48 hours and 10 days after death, and adrenal gland apoplexy was detected in all the cases. Microbiological examinations identified *Neisseria meningitidis*, which was accountable for Waterhouse–Friderichsen syndrome. Diplococci were isolated from three cadavers that underwent forensic dissection between 2 and 3 days after death. The remaining two cadavers showed polymicrobial contamination, and a polymerase chain reaction technique was necessary to identify the pathogen. We assumed that the microbial overlap could lead to diagnostic mistakes and conceal the identification of the lethal pathogen. Therefore, we suggest using molecular techniques for a postmortem interval (PMI) longer than 72 hours. Classical microbiological examination should be performed for PMI within 72 hours.

Keywords

Autopsy; Forensic Microbiology; Waterhouse-Friderichsen Syndrome; Post-Mortem Interval.

INTRODUCTION

Waterhouse–Friderichsen syndrome (WFS) is a catastrophic sepsis¹ with a higher incidence in children,²⁻⁴ and is usually associated with adrenal gland hemorrhage, which is considered-along with the disseminated intravascular coagulation (DIC)-as the most fearsome complication. Due to the fact that symptoms may be non-specific, the *in-vivo* diagnosis of adrenal gland hemorrhage is difficult to define⁵ and is usually identified at autopsy.¹ Despite their protected position in the retroperitoneal space (an area rich in adipose-connective tissue),⁶ at autopsy examination hemorrhage of the adrenal glands can be found in several conditions as traumatism,⁷ or rare cases of infection-related⁸ sudden death,⁹ such as WFS. People suffering from WFS may initially show unspecific symptoms, such as fever—with a temperature always

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higher than 38°C—then, suddenly, the general conditions worsen and progressive hypotension occurs along with shock and disseminated intravascular coagulation. In 50%-60% of cases² cutaneous purpura may be observed. The clinical development of the disease is variable; it may be: (i) mild and undergo spontaneous resolution; (ii) sub-acute with an insidious and slow development; or (iii) fulminant and lead to death in a short period of time. The latter event usually occurs after the setting of the first symptoms, and these episodes-with or without involvement of the meninges—are complicated by the septic process, the disseminated intravascular disease, and massive hemorrhage of the adrenal glands (distinct WFS). There are three fundamental elements necessary for the identification of this lethal form:¹⁰

- 1. Etiological factors (meningococcal hematogenous dissemination or fulminant pneumococcal hematogenous dissemination)
- 2. Clinical presentation (severe and rapid progression of the infectious setting)
- 3. Peculiar anatomical lesions (cerebral-spinal meningitis signs with compromised neural-axons and bilateral adrenal gland hemorrhage (cases of unilateral adrenal gland hemorrhage have been described in literature).¹¹

The mortality rates of the meningococcal/ pneumococcal disease, not associated with WFS, are in the range of 10%-30%,³ sensibly increasing to 95% in cases where WFS is associated. The aim of this study is to (i) verify the sensitivity of post-mortem microbial examination on victims deceased due to certified infective pathologies (WFS) who undergo forensic dissection at different periods of time; and (ii) try to delineate the time interval in which it is still possible to microbiologically detect *Neisseria meningitidis*.

CASE REPORTS

Case 1

A 2-month-old male baby was carried to Accident & Emergency (A&E) with persistent fever. After clinical examination, the patient was prescribed with antipyretics (paracetamol), but the same day he developed spreading petechiae. The medical staff requested cerebrospinal fluid bacterial exams. Unfortunately, the little patient died before the results of the exams became available. An autopsy was performed 10 days after death. The cadaver had several cutaneous petechiae and adrenal glands hemorrhage. The microbiological post-mortem exams showed an autolytic multi-bacterial proliferation on leptomeninges and cardiac blood. A molecular investigation (polymerase chain reaction [PCR]) on the same tissue and blood resulted positive for *N. meningitidis*. Histological examinations were not allowed by the Judicial Authority.

Case 2

A 5-month-old male baby was carried to A&E with fever. The medical staff prescribed an antipyretic (paracetamol). The following day, the baby became cyanotic and developed a cutaneous exanthema. The baby died while clinical exams were still being carried out. The autopsy was performed 3 days after death. The cadaver had several cutaneous petechiae, congestion of the leptomeninges with opaque coloration, deposits of fibrin, purulent exudate of yellow-greenish color on the convexities of the hemispheres, and bilateral apoplexy of the adrenal glands. Post-mortem microbial exams revealed the presence of *N. meningitidis* on cerebral swabbing. Histological examinations of brain, cerebellum, and pons showed a pronounced edema of the leptomeninges, inflammatory granulocyte infiltration, few lymphocytes, and marked dilatation and hematic stasis of the leptomeningeal vascular system. Histological examinations of the adrenal glands showed acute hemorrhagic infiltration of both glands.

Case 3

A 3-year-old girl was carried to A&E presenting petechiae, vomiting, and fever. The physicians suspected meningitis and administered an antibiotic (ceftriaxone), but she suddenly died. The autopsy examination was performed 4 days later. The cadaver had widespread cutaneous petechiae, necrotic hemorrhage of adrenal glands, and purulent exudate in the frontal-parietal lobes and cerebellum. Postmortem microbiological exams revealed Gram-negative bacterial cells on sampled purulent areas. The molecular investigations (PCR) were positive for *N. meningitidis*. Histological examinations showed dilated and marked leukostasis

(formed by neutrophils) in the leptomeninges, widening of perivascular and pericellular spaces (due to severe cellular edema), adrenal glands apoplexy, septal microbial micro-abscesses, and marked edematous dissociation of cardiac fibers and pericardial connective tissue.

Case 4

A 17-year-old boy was taken to A&E with fever (39.5 °C) and sore throat. The physicians suspected meningitis and administered an antibiotic (ceftriaxone). The boy died after 1 day. The forensic examination was performed 3 days after death. The corpse had opaque leptomeninges at cerebral convexities. The adrenal glands showed areas with necrotic-hemorrhagic aspect. Postmortem microbial investigations revealed the presence of *N. meningitidis* on cardiac blood and brain samples. Histological examinations revealed early signs of meningitis on the leptomeninges, and hemorrhagic infiltration of both adrenal glands.

Case 5

A 23-year-old female presented with fever, tremor, pain, and developed paresthesia to the limbs 24 hours after the onset of the first symptoms. She was taken to one of the main hospitals of the city of Milan where she died minutes after admission. The autopsy examination was performed 5 days after death. The corpse showed diffuse cutaneous petechiae, which was also detected at dissection on the temporal muscles, brain, esophagus, pleurae, pericardium, stomach, intestines, and liver. The adrenal glands showed bilateral hemorrhagic infiltration. Postmortem microbiological cultures of viscera and fluids were negative. The PCR was positive for *N. meningitidis*. The histological examinations showed a multi-organ hemorrhagic petechial syndrome, a perivascular sleeve of neutrophils in an intra-parenchymal brain vessel with an initial infiltration of the nearby parenchyma (mild stage meningitis), bilateral adrenal gland hemorrhage, and acute multi-focal bacterial myocarditis.

In all cases, the cause of death was identified in WFS caused by *N. meningitidis.* (Table 1).

Autopsy presentation

Diagnosis of WFS can be arduous: cadavers coming to the attention of the forensic practitioner with early symptoms and with a lack of cutaneous macroscopic signs, can pose difficulties in terms of differential diagnosis between WFS and a milder viral infection. On the other hand, in cadavers showing an *intra-vitam* history of high fever, spreading cutaneous exanthema, and progressive shock, WFS must always be suspected.² In these cases, autopsy examinations request the adoption of adequate measures of protection. In a paper edited by the Centers for Disease

Table 1. Demographic data, autopsy and microbiological results of the 5 cases of Waterhouse-Friderichsen Syndrome

Case	Sex	Age	Autopsy sampling	Skin	Brain	Adrenal apoplexy	Other organs' signs	Agent	Cause of death
1	Μ	2 m	Meninges, cardiac blood	WP		yes		<i>N. meningitidis</i> (with PCR)	WFS
2	Μ	5 m	cerebral swabs	WP	opaque meninges, congestion, purulent deposits	yes		N. meningitidis	WFS
3	F	3 у	purulent material	WP	opaque meninges, purulent deposits	yes	endocardial petechiae, bacterial involvement of the myocardium	<i>N. meningitidis</i> (with PCR)	WFS
4	М	17 y	brain and cardiac blood		opaque meninges	yes		N. meningitidis	WFS
5	F	23 у	fluids and organs	WP	petechiae in the brain, cerebellum and temporal muscles	yes	Visceral petechiae, bacterial involvement of the myocardium, renal micro - thrombi	<i>N. meningitidis</i> (with PCR)	WFS

F = feminine; M = masculine; m = month; PCR = polymerase chain reaction; y = year; WFS = Waterhouse Friderichsen Syndrome; WP = Widespread Petechiae.

Control and Prevention (CDC),¹² since the pathology is contagious and airborne, the preventive measures recommended for the microbiological safety of the laboratory personnel potentially exposed to aerosolized *N. meningitidis* should be vaccines; however, nothing is stated about preventive measures for mortuary staff.

When WFS is full-blown, it can be detected macroscopically at the external examination thanks to characteristic signs, such as cutaneous petechiae (Figure 1A), and at internal dissection due to purulent meningeal and cerebral matter and adrenal gland hemorrhage —small bleedings usually located at the cortical-medullary junction that can be detected with a careful histological examination.⁶ The sampling of these cadavers requests a befitting selection of the biological fluids that must be collected with sterile syringes and stored in sealed vials: cardiac blood^{9,13} and cerebrospinal fluid^{3,14} are the most fitting samples; moreover, the examiner can proceed with the sampling of cadaveric parenchyma with sterile lancet and clamps, and storage in sealed vials. It is also advisable to sample the spleen,^{9,14} heart, and liver.¹⁵ In addition, an extensive cerebral sampling is recommended, including: leptomeninges,³ cerebral hemispheres' base and convexity (frontal, temporal, parietal, and occipital lobes), cerebellum (hemispheres and vermis), brainstem (midbrain, pons, and medulla oblongata), and spinal cord (cervical, thoracic, abdominal, and sacrum-coccygeal). The CDC guidelines¹² recommend to start a post-exposure antimicrobial chemoprophylaxis within 24 hours.

This study was conducted at the Legal Medicine Section of the University of Milan. We selected five

patients (three males and two females aged between 2 months and 23 years) who suddenly died after showing symptoms of persistent fever. The autopsy was performed on all five patients between 48 hours and 10 days after death. Three cases (2, 3, 4) had adrenal apoplexy (Figure 1B) and meningitis signs (Figure 1C). Two cases (1, 5) did not show signs of meningitis.

Laboratory Procedures

During autopsy, a certain amount of biological fluids was destined for microbiological examination with a thin smear on at least two microscope slides. One of these slides was dyed with Gram cytological staining in order to highlight the presence of possible bacterial cells while the other slide was dyed with cytological hematoxylin and eosin (H&E) in order to evaluate the possible presence of inflammatory infiltrate. The remaining samples were addressed to standard cultural exams. If any cultures were negative or characterized by postmortem autolytic growth, it was considered necessary to continue with higher level procedures, such as molecular tests.

In order to proceed with standard histopathological analysis, slides had to be stained with H&E, histologic Gram staining, and phosphotungstic acid hematoxylin. After the staining process, fibrin clots were detected (Figure 2A) along with neutrophil infiltration in cerebral parenchyma (Figure 2B), cortical hemorrhage of the adrenal glands (Figure 2C), and multifocal acute bacterial myocarditis with several neutrophil infiltrations (Figure 2D).

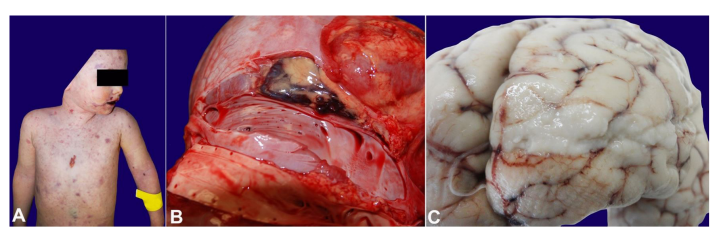


Figure 1. (Case 3) $- \mathbf{A} - \mathbf{G}$ ross examination of the corpse's body showing diffuse cutaneous petechiae; **B** and **C** - opacification of the meninges with deposition of purulent material over the frontal and parietal lobes, vermis and cerebellar lobes.

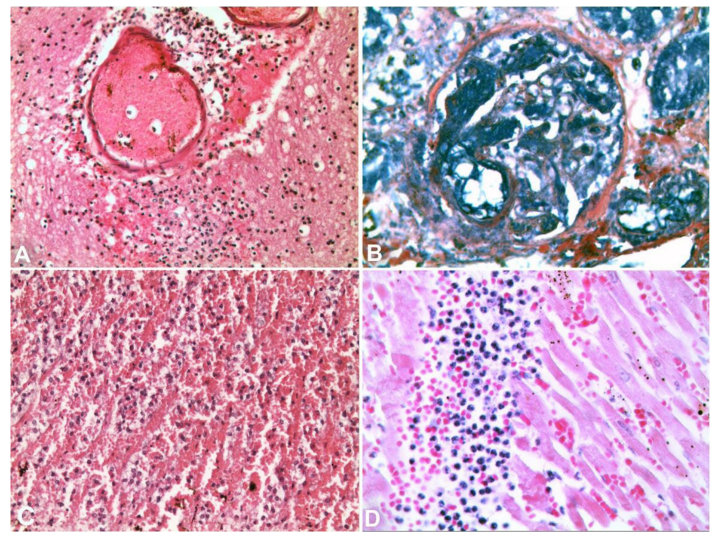


Figure 2. Photomicrographs of: **A** – Brain - small intra-parenchymal cerebral vessel with perivascular sleeve of neutrophils and surrounding parenchymal infiltration (H&E, 200X); **B** – Kidney – glomerular loops with necrotic micro-foci of the capillary vessels congruent with deposits of fibrinoid material, expression of an intravascular disseminated coagulation (PTAH, 400X); **C** – adrenal gland - (H&E, 200X)- massive cortical hemorrhage of the adrenal glands with adrenal cells remnants; **D** – Heart - diffuse multifocal acute bacterial myocarditis with several neutrophils infiltrations associated to intramuscular micro-hemorrhage and fibers' necrosis (H&E 400X).

A recent acquisition in Forensic Sciences is the assessing of molecular and genetic amplification. This procedure, using a specific gene primer, enables the detection of CrgA—an adhesion regulatory protein responsible for the adhesion of *N. meningitidis* to its target cells,⁶—in cerebrospinal fluid, cerebral parenchyma with leptomeninges (especially the arachnoid mater), spleen tissue,^{14,15} and cutaneous purpura lesions.¹⁶ International studies¹ use the serum concentration of procalcitonin as a postmortem biochemical marker of sepsis. Normal blood levels of 0.5 ng/mL, or lower, can rise to 10 ng/mL in patients deceased due to sepsis; recently, vitreous humor has also been considered as a useful substrate to isolate

N. meningitidis, since this humor is resistant to blood biochemical postmortem changes.¹⁷

Etiology

The etiological agent often remains unidentified due to intrinsic difficulties related to the characteristics of the analyzed tissues and to its peculiar features:¹⁴ the etiological prevalence, in more than 80%¹⁸ of cases, is generally supported by *N. meningitidis*, a Gram-negative coccus. In preparations with Gram staining, this coccus appears as a kidney-like shape, with cells paired up through the flattened side and with a "coffee-bean-like" aspect. This bacterium can be intracellular or extracellular (Figure 3), is often

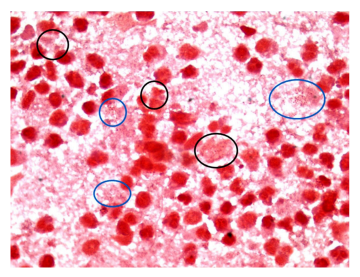


Figure 3. Cerebral smear with Gram staining highlighting the diagnostic difficulties on a substrate of granulocytes neutrophils undergoing the autolytic process, but still recognizable. Evidence of Gram-negative diplococci with "coffee bean-like" aspect, isolated or paired in couples or tetrads, organized both in the cellular cytoplasm of the neutrophils (black circles) and outside the cytoplasm (blue circles).

arranged in couples (diplococcus), and is found in the nasopharynx of 3%-30% of healthy individuals,¹⁵ who can be considered as asymptomatic carriers but can develop meningitis during or after flu infections.¹⁹ Other virulent organisms like Klebsiella,²⁰ Ewingella,⁹ Haemophilus influenzae,²¹ Streptococcus pyogenes,²² Streptococcus pneumonia,^{7,22} and Pasteurella²³ occasionally can be detected as etiological agents of WFS and isolated in cadaveric samples. The etiological agent was not immediately identified with standard microbiological techniques in three cases (1, 3, 5) with a postmortem interval (PMI) longer than 72 hours; instead, the bacterium was identified with real-time PCR. In the remaining cases (2, 4) with a PMI within 72 hours, N. meningitidis was identified with routine microbiological cultures.

DISCUSSION

We wanted to verify the sensitivity of the postmortem microbial testing on a group of five patients who died due to WFS and underwent forensic dissections at different times—PMI between 2 and 10 days—to try to evaluate the interval time in which it was still possible to microbiologically detect the bacterial agent. We can draw the following conclusions:

- 1. The PMI should be less than 72 hours in order to perform a reliable microbiological examination on cadaveric samples: in the cases with PMI greater than 72 hours (1, 3, 5) it was not possible to isolate and detect the lethal microbial agent; therefore, real-time PCR was necessary. The other two cadavers (2, 4) were stored for less than 72 hours in refrigerating rooms. We were able to investigate both with cultures of cadaveric matter and with cerebral sampling stained with cytological Gram. These two methods were simpler than the molecular ones. We concluded that there is a multi-microbial overlap due to a PMI longer than 72 hours. This can lead to diagnostic mistakes and conceal the identity of the pathogen involved in the mechanisms of death.²⁴ Therefore, it is essential to use a molecular bacterial method in all cases of suspected WFS with a wide postmortem interval.
- Leptomeninges are targeted by meningococci in WFS cases even if they are not macroscopically involved. In fact, the leptomeninges were clean at the macroscopic examination of cases 4 and 5, but the histological examination revealed a variable stage—mild to moderate,³—of meningitis.
- 3. The septic invasion from meningococci, as in cases 3 and 5, must be validated by histological exam of samples taken from different organs, especially myocardium. Bacterial myocarditis is commonly observed because of the septic processes. Therefore, the actual cause of death may be due to shock, DIC, or myocarditis³ in such cases.

CONCLUSION

WFS is characterized by high mortality, sudden onset of symptoms, rapid clinical development, and unexpected death. These factors open various medical-legal issues related to the cause of death, the search for sources of infection, and the assessment of any medical malpractice. What must be stated is that WFS has to be distinguished from sudden infant death syndrome (SIDS), since autopsy results always provide proof of the cause of death, while macroscopic and microscopic investigations in cases of SIDS are negative to the assessment of a precise cause of death.³ Moreover, in terms of medical malpractice, the answers requested by forensic investigations are whether or not there was a missed or delayed diagnosis/treatment, which may have prevented lethal outcomes.⁴ Victims' relatives, especially when children are involved, often blame the general practitioner or pediatrician who took the little patient for the first clinical observations; however, in these patients, it is not easy to discern such a lethal infection from common flu or enteritis,³ since symptoms such as fever or inflammation of the upper airways may be commonly expressed.⁴ When the diagnosis of WFS becomes clear, its development cannot be stopped, and there is no effective therapy for it.

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