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Death after Quadrivalent Human Papillomavirus (HPV) Vaccination: Causal or Coincidental?
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Abstract

**Background:** The proper understanding of a true risk from vaccines is crucial for avoiding unnecessary adverse reactions (ADRs). However, to this date no solid tests or criteria have been established to determine whether adverse events are causally linked to vaccinations.

**Objectives:** This research was carried out to determine whether or not some serious autoimmune and neurological ADRs following HPV vaccination are causal or merely coincidental and to validate a biomarker-based immunohistochemical (IHC) protocol for assessing causality in case of vaccination-suspected serious adverse neurological outcomes.

**Methods:** Post-mortem brain tissue specimens from two young women who suffered from cerebral vasculitis-type symptoms following vaccination with the HPV vaccine Gardasil were analysed by IHC for various immunoinflammatory markers. Brain sections were also stained for antibodies recognizing HPV-16L1 and HPV-18L1 antigen which are present in Gardasil.

**Results:** In both cases, the autopsy revealed no anatomical, microbiological nor toxicological findings that might have explained the death of the individuals. In contrast, our IHC analysis showed evidence of an autoimmune vasculitis potentially triggered by the cross-reactive HPV-16L1 antibodies binding to the wall of cerebral blood vessels in all examined brain samples. We also detected the presence of HPV-16L1 particles within the cerebral vasculature with some HPV-16L1 particles adhering to the blood vessel walls. HPV-18L1 antibodies did not bind to cerebral blood vessels nor any other neural tissues. IHC also showed increased T-cell signalling and marked activation of the classical antibody-dependent complement pathway in cerebral vascular tissues from both cases. This pattern of complement activation in the absence of an active brain infection indicates an abnormal triggering of the immune response in which the immune attack is directed towards self-tissue.

**Conclusions:** Our study suggests that HPV vaccines containing HPV-16L1 antigens pose an inherent risk for triggering potentially fatal autoimmune vasculopathies.

**Practice implications:** Cerebral vasculitis is a serious disease which typically results in fatal outcomes when undiagnosed and left untreated. The fact that many of the symptoms reported to vaccine safety surveillance databases following HPV vaccination are indicative of cerebral vasculitis, but are unrecognized as such (i.e., intense persistent migraines, syncope, seizures, tremors and tingling, myalgia, locomotor abnormalities, psychotic symptoms and cognitive deficits), is a serious concern in light of the present findings. It thus appears that in some cases vaccination may be the triggering factor of fatal autoimmune/neurological events. Physicians should be aware of this association.

Keywords: HPV vaccines; Serious adverse reactions; Cerebral vasculitis; Vasculopathy; Autoimmunity; Molecular mimicry; Immune complexes; Autoantibodies

Introduction

In the past several decades, there have been numerous studies and case reports documenting neurological and autoimmune adverse reactions (ADRs) following the use of various vaccines. Arthritis, vasculitis, systemic lupus erythematosus (SLE), encephalopathy, neuropathy, seizure disorders and autoimmuno demyelinating disease syndromes are the most frequently reported serious adverse events [1-13]. Although a clear temporal relationship between the administration of a vaccine and the adverse event is sometimes observed, in the vast majority of cases no causal connection can be demonstrated. Thus, it is often concluded that, (i) the majority of serious ADRs that occur post-vaccination are coincidental and unrelated to the vaccine [14] and, (ii) true serious vaccine-related ADRs (i.e., permanent disability and death) are extremely rare [15]. There are however several important reasons why causality is rarely established with regard to vaccination-associated ADRs. These include: the criteria for causality are poorly defined [6,16,17]; the latency period between vaccination and autoimmunity can range from days to years (individuals’ susceptibility factors most likely playing a role in determining the temporal onset, time course, and severity of symptoms) [6]; neurological outcomes, as in other neurological disorders may take considerable periods to manifest as obvious pathology [18]; post-vaccination adverse manifestations can be atypical and might not be compatible with a defined autoimmune or neurological disease [6]; individual susceptibility factors are not considered and a “one-size fits all” principle is assumed [19]; a triggering role of the vaccine in the adverse outcome is not considered [20].

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Quadrivalent human papillomavirus (qHPV) vaccine Gardasil was licensed in 2006 by the U.S. Food and Drug Administration (FDA) following a fast track approval process [21]. In pre- and post-licensure epidemiological studies no autoimmune safety concerns associated with Gardasil vaccination were identified [22,23] and the vaccine has been considered to have a remarkably good safety profile [24]. However, a careful scrutiny of Gardasil safety trials shows evidence of significant flaws in study design, data reporting and interpretation [25-28]. Irrespective of these latter observations, it is important to note that epidemiological studies only test for “association” and not “causation”, thus providing unreliable estimates of true risks. On the other hand, data from numerous case reports documenting serious autoimmune and neurological complications following HPV vaccination continue to raise concerns [11,12,29-35]. Nonetheless, the precise etiology of these post-HPV vaccination associated phenomena has been elusive and hence causality remains unascertained.

We have recently developed an immunohistochemical (IHC) protocol based on analyzing two cases of sudden and unexplained death following vaccination with the qHPV vaccine Gardasil. This protocol has been developed for the purpose of determining whether the serious autoimmune and neurological manifestations reported following HPV vaccination are causal or merely coincidental.

Patients and Methods

Case 1

A 14-year-old female with a previous history of migraines and oral contraceptive use developed more severe migraines, speech problems, dizziness, weakness, inability to walk, depressed consciousness, confusion, amnesia and vomiting 14 days after receiving her first qHPV injection. These symptoms gradually resolved. However, a full autopsy no major abnormality was found anatomically, microbiologically or toxicologically that might have been regarded as a potential cause of death. Histological analysis of the brain hippocampus, cerebellum, and watershed cortex allegedly revealed no evidence of neuronal loss or neuroinflammatory changes. However, the autopsy report did not specify which immune antibodies and stains were used for histological investigations.

Case 2

A 19-year-old female without a relevant medical history and taking no drugs expired in her sleep, approximately 6 months after her third and final qHPV vaccine booster and following exacerbation of initial vaccination-related symptoms. She had last been seen alive by her parents the previous evening. Her symptoms started after the first qHPV injection when she developed warts on her hand that persisted throughout the vaccination period. In addition, she suffered from unexplained fatigue, muscle weakness, tachycardia, chest pain, tingling in extremities, irritability, mental confusion and periods of amnesia (memory lapses). The autopsy was unrewarding and failed to determine the exact cause of death. Internal examination revealed some minor changes involving the gallbladder and the uterine cervix (both of which on further examination by microbiological studies and histology revealed no significant disease). After a full autopsy no major abnormality was found anatomically, microbiologically or toxicologically that might have been regarded as a potential cause of death. Histological analysis of the brain hippocampus, cerebellum and watershed cortex allegedly revealed no evidence of neuronal loss or neuroinflammatory changes. However, the autopsy report did not specify which immune antibodies and stains were used for histological investigations.

Tissue samples

Paraffin-embedded brain tissue specimens collected at autopsy from the two cases described above were used in this study. The tissues analyzed by IHC included the cerebellum, hippocampus, choroid plexus and watershed cortex (Case 1), and cerebellum, hippocampus, choroid plexus, portions of the brainstem (medulla, midbrain, pons), right basal ganglia, right parietal and left frontal lobes (Case 2). Sections were cut at 5 μm and mounted on glass slides.

IHC procedures

The paraffin brain sections were dewaxed, rehydrated and incubated for 15 min in methanol containing 3% H₂O₂ to block endogenous peroxidase activity. The sections were then incubated in 1% saponin in 1xPBS for 1 hr at room temperature (RT) and subsequently pre-treated by boiling for 20 min in an appropriate antigen retrieval solution (Table 1) to facilitate antigen retrieval and to increase membrane permeability to primary antibodies (Abs). The sections were then blocked in 1xPBS containing 0.1% Tween and 10% normal goat serum for 15 min in methanol containing 3% H₂O₂. The sections were incubated in 1% saponin in 1xPBS for 1 hr at RT. Following the blocking step, the sections were incubated with appropriate dilutions of the primary Ab in the blocking solution (Table 1) overnight at 4°C in a humidified chamber.

Subsequently, sections were rinsed three times in 1xPBS and incubated with biotinylated, affinity-purified anti-immunoglobulin G (IgG) secondary Ab at 1:200 dilution (Vector Labs, Inc), and then with the avidin-biotin complex (ABC)-immunoperoxidase Vectastain Elite ABC kit (Vector Laboratories, Burlingame, Calif). The positive reaction was visualized by 3,3-diaminobenzidine (DAB) peroxidase according to standard methods. The sections were counterstained with methyl green, dehydrated, coverslipped, and observed on a Zeiss Axiosvert microscope (Carl Zeiss Canada Ltd., Toronto, ON) connected to a computerized system with a photo camera. Images at 10x and 40x magnification were captured using AxiosVision 4.3 software.

Results and Discussion

Principal findings

The results from our IHC examinations of brain tissue specimens from two young women who died following vaccination with the
The qHPV vaccine Gardasil showed strong evidence of an autoimmune vasculitis triggered by the cross-reactive HPV-16L1 antibodies binding to the wall of cerebral blood vessels (Figures 1 and 2). In addition, there was clear evidence of the presence of HPV-16L1 particles within the cerebral vasculature with some HPV-16L1 particles adhering to the blood vessel walls (Figures 1C, 2C and 2D). In contrast, HPV-18L1 antibodies did not bind to cerebral blood vessels nor any other neural tissues (Figure 1D).

Histopathologically, immune-mediated vasculitis is typified by excessive recruitment and vascular adhesion of T lymphocytes, increased complement and MHC II signaling and complement-dependent deposition of immunoglobulin G–immune complexes to cerebral vasculature [5,36-40]. Increased expression of proteolytic matrix metalloproteinases (MMPs) also characterizes immune vasculopathic syndromes [41-43]. The MMPs play a major role in both the progression of inflammatory infiltrates and vessel destruction [41-43]. The ensuing vascular damage is manifested in hemorrhagic and ischemic brain tissue lesions. Notably, there was clear and consistent evidence of all these pathogenic immune processes in the brain tissue specimens from both young women (Figures 3-8).

The finding of a cerebral edema following autopsy in Case 2 is strongly suggestive of a focal blood-brain barrier breakdown [10,45]. Additionally, H&E staining showed clear evidence of hemorrhages (Figure 8), which were also present in Case 1 (Figure 7). Disruption of the blood-brain barrier manifesting in hemorrhagic tissue lesions could have resulted from both vasculitis and deleterious effects of excessive levels of glia-derived inflammatory cytokines. Although the autopsy failed to show evidence of microglial and inflammatory reactions in both cases, this was likely because no glia-specific markers were used in the histopathological analyses of brain tissue specimens. In contrast, results from our IHC analysis using micro- and astroglia specific markers (Table 1) showed exceptionally intense micro- and astrogliosis in all brain tissue sections examined from both Case 1 and Case 2 (Figures 9 and 10). Microglia are the brain’s resident immune cells and their excessive activation can lead to irreversible neurodestructive and pro-inflammatory processes in the brain [46,47].

It is also well known that activated microglia increase the permeability of the blood-brain barrier to other inflammatory factors and to trafficking lymphocytes [48]. Moreover, microglial aggregation in the brain is also recognized as a marker for hypoxic-ischemic brain injury [49], the latter diagnosed by the coroner in Case 2. Both microglia and astroglia are activated by a variety of immune insults, including aluminum vaccine adjuvants which are present in both HPV vaccines Gardasil and Cervarix. In addition, animal experiments show that only the latter applies in both cases described.

The table below lists the primary antibodies used in the study:

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Type</th>
<th>Epitope/Specificity</th>
<th>Pre-treatment/antigen retrieval</th>
<th>Dilution</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1q</td>
<td>Monoclonal, Anti-human</td>
<td>Complement, classical pathway</td>
<td>0.25 mM EDTA buffer pH 9.0</td>
<td>1:100</td>
<td>Abcam</td>
</tr>
<tr>
<td>C5b-9</td>
<td>Monoclonal, Anti-human</td>
<td>Complement, membrane attack complex</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Abcam</td>
</tr>
<tr>
<td>IgG, IgM, IgA</td>
<td>Polyclonal, Anti-human</td>
<td>Immunoglobulin complexes</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Dako</td>
</tr>
<tr>
<td>HPV-16L1</td>
<td>Monoclonal, Anti-HPV-16</td>
<td>HPV-16 major capsid protein L1</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Abcam</td>
</tr>
<tr>
<td>HPV-18L1</td>
<td>Monoclonal, Anti-HPV-18</td>
<td>HPV-18 major capsid protein L1</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Abcam</td>
</tr>
<tr>
<td>HLA-DR (MHC II)</td>
<td>Monoclonal, Anti-human</td>
<td>Major histocompatibility complex class II</td>
<td>0.25 mM EDTA buffer pH 9.0</td>
<td>1:100</td>
<td>Dako</td>
</tr>
<tr>
<td>CD3</td>
<td>Monoclonal, Anti-human</td>
<td>T-cell lymphocytes</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Abcam</td>
</tr>
<tr>
<td>MMP-9</td>
<td>Polyclonal, Anti-human</td>
<td>Matrix metalloproteinase 9</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:100</td>
<td>Calbiochem</td>
</tr>
<tr>
<td>GFAP</td>
<td>Monoclonal, Anti-mouse/human</td>
<td>Reactive astrocytes</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:500</td>
<td>Wako</td>
</tr>
<tr>
<td>Iba-1</td>
<td>Polyclonal, Anti-mouse/human</td>
<td>Activated microglia</td>
<td>0.1 M Citric Acid buffer pH 6.0</td>
<td>1:500</td>
<td>Wako</td>
</tr>
</tbody>
</table>

**Table 1:** Panel of primary antibodies used in the study.
Vascular immunostaining with anti-HPV-16L1 (A-C) and anti-HPV-18L1 antibodies (D). Vascular wall of two small blood vessels in the watershed cortex showing positive immunoreactivity for virus-like particles (VLPs) of the recombinant major capsid (L1) protein of HPV-16 (A, B). Note that some HPV-16L1 particles tissue have infiltrated the brain tissue parenchyma adjacent to the vessel (B, arrow). Intravascular accumulation of HPV-16L1 VLPs in the watershed cortex (C). Note that some HPV-16L1+ cells are adhering to the endothelial lining of the vessel wall (C, arrows). Lack of HPV-18L1 immunoreactivity in the watershed cortex (D). HPV-18L1 immunoreactivity was also absent in other brain tissue specimens from both Case 1 and Case 2 (not shown). Magnification 40x.

Vasculitis has long been recognized as a possible severe ADR to vaccination [5,54-57]. For example, Carvalho and Shoenfeld described a case of polyarteritis nodosa, a rare, life-threatening, necrotizing vasculitis that affects medium-sized arteries following the administration of the hepatitis B vaccine in a 14-year-old boy who had no relevant previous history and who was not taking any drugs [5]. There have been numerous other case reports of vasculitis post-hepatitis B vaccination. In IHC examinations of these cases, no virus B antigen was detected in the blood vessels and no proof of cause and effect was established [55,57]. However, the possibility that the vaccine-derived hepatitis B antigen might have caused autoimmunity by molecular mimicry was not investigated. In fact, to the best of our knowledge, our study is the first to show direct evidence of a vaccine antigen-induced autoimmune vasculitis in the human central nervous system (CNS).

It is worth emphasizing that molecular mimicry (whereby the vaccine antigen resembles a host antigen) is generally accepted as a mechanism by which vaccines can trigger autoimmune diseases. Thus, inflammation of various types, vasculitis may underlie diverse diseases, making its diagnosis difficult [37,39]. Vasculitis can affect blood vessels including arteries and veins of all sizes in all areas of the body, resulting in a variety of clinical neurological manifestations [37,39].

The brain is particularly sensitive to ischemia. Vasculitis of the nervous system is thus of paramount importance to clinicians as it almost inevitably leads to permanent injury and disability when unrecognized and left untreated [37]. Cerebral vasculitis is thought to be rare [40]. However, because clinical presentation of cerebral vasculitis is highly variable with fluctuating signs and symptoms [40], and because it frequently underlies many diverse inflammatory diseases (i.e., SLE [41] and bacterial meningitis [43]), it is probable that many cases of cerebral vasculitis remain undiagnosed and/or misdiagnosed.

Typical symptoms of cerebral vasculitis include severe headaches, orthostatic dizziness, syncope, seizures, tremors and tingling, weakness, locomotor deficits, cognitive and language impairments [39,51,52]. Note that the vast majority of these symptoms were experienced by both Case 1 and Case 2 post-qHPV vaccination (Table 2).

Vasculitis has long been recognized as a possible severe ADR to vaccination [5,54-57]. For example, Carvalho and Shoenfeld described a case of polyarteritis nodosa, a rare, life-threatening, necrotizing vasculitis that affects medium-sized arteries following the administration of the hepatitis B vaccine in a 14-year-old boy who had no relevant previous history and who was not taking any drugs [5]. There have been numerous other case reports of vasculitis post-hepatitis B vaccination. In IHC examinations of these cases, no virus B antigen was detected in the blood vessels and no proof of cause and effect was established [55,57]. However, the possibility that the vaccine-derived hepatitis B antigen might have caused autoimmunity by molecular mimicry was not investigated. In fact, to the best of our knowledge, our study is the first to show direct evidence of a vaccine antigen-induced autoimmune vasculitis in the human central nervous system (CNS).

It is worth emphasizing that molecular mimicry (whereby the vaccine antigen resembles a host antigen) is generally accepted as a mechanism by which vaccines can trigger autoimmune diseases. Thus,
antibodies and T cells that are produced to destroy the vaccine antigen also attack structurally similar self-antigens in different tissues (i.e., the wall of blood vessels) [5,6,35]. The fact that vaccination is often intended to prevent a disease and thus carried out in the absence of an active infection in the host, implies that the risk of autoimmunity may be exacerbated if there is structural similarity between the vaccine antigen and the host tissue. The reason for this is two-fold. Firstly, vaccination produces a much higher and sustained level of antibodies compared to natural infection (i.e., Gardasil-induced HPV-16 antibody titers are 10-fold higher than natural HPV infection titers [58]). Secondly, in the absence of an actual infectious agent (i.e., HPV-16 virus), the vaccine-induced antibodies are likely to preferentially bind to host antigens.

**Vasculitis and HPV vaccination**

Recently two cases of vasculitis affecting young teenage girls following HPV vaccination have been reported. In both cases there was no history of preceding infection, and the strong temporal relationship between the administration of the bivalent HPV vaccine (Cervarix) and the development of vasculitis was notable [32].

Our search of the Vaccine Adverse Event Reporting System (VAERS) internet database [59] revealed numerous reports of post-HPV vaccination-associated vasculitis. An analysis of these cases shows that post-vaccination vasculitis-related symptoms most typically manifest within the first three to four months of vaccination (Table 3), as was also reported in Case 1 and Case 2 described herein. Moreover, we noted a striking similarity between the vasculitis-related symptoms reported to VAERS and those experienced by Case 1 and Case 2 (Table 2). Although a report to a passive vaccine safety surveillance system such as VAERS does not by itself prove that the vaccine caused an ADR, it should be noted that many VAERS reports also include detailed records of diagnostic laboratory analyses and clinician’s follow-ups and their expert diagnosis. Notably, some of these reports include medically confirmed cases where the diagnosis of immune-mediated vasculitis was ascertained (i.e., VAERS ID# 425345-1, 436679-1; Table 3).

The precise etiology and the role of HPV vaccination in vasculitis cases reported to VAERS remained undetermined. However, we note that the histopathological examinations when conducted were very limited in scope (Table 3). Specifically, the possibility of HPV vaccine-induced autoimmunity via molecular mimicry due to cross-reactivity between vaccine antigens and host vascular structures was neither investigated nor considered. The reason for such omissions in histopathological analyses of vaccine-suspected autoimmune pathologies is unclear, especially since medical scientists generally

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**Figure 5:** Case 1: Vascular (arrows) and perivascular (arrowheads) accumulation of HPV-16L1 (A,B), MHC class II (C,D), CD3 (E,F), C5b-9 (G,H) and IgG-IgM-IgA (I,J) positive cells. WCx=watershed cortex; Hippo=hippocampus; Crb=cerebellum. Magnification 10x (A) and 40x (B-J).

**Figure 6:** Case 2: Vascular accumulation of HPV-16L1 (A-C,M), MHC class II (D-F), CD3 (G-I), C5b-9 (N) and IgG-IgM-IgA (O) positive cells. Note the consistent pattern of HPV-16L1, MHC II, CD3 and C1q immunostaining (A-L), where majority of immunopositive cells are found adhering to the walls of cerebral blood vessels (arrows). Some MHC II+ cells (D, arrowhead) and CD3+ T cells are also found in the perivascular rim (G, arrowhead) and some C1q+ cells are infiltrating deep into the brain parenchyma adjacent to the vessel (K, arrowheads). Note the dense accumulation of HPV-16L1 (M), C5b-9 (N) and IgG-IgM-IgA (O) immunopositive cells in the cerebellar blood vessels. Hippo=hippocampus; Crb=cerebellum; Midb=midbrain; RBas=right basal ganglia. Magnification 40x.

**Figure 7:** Case 1. Hematoxylin & Eosin (H&E) stain of blood vessels in the cerebellum (A), and the watershed cortex (B,C) showing hemorrhages. Note the presence of red blood cells in the brain tissue parenchyma adjacent to the blood vessels indicating brain tissue hemorrhage (arrows). Magnification 40x.

**Figure 8:** Case 2. Hematoxylin & Eosin (H&E) stain of arteries in the medulla (A), hippocampus (B) and the right basal ganglia (C) showing hemorrhages. Note the presence of red blood cells in the brain tissue parenchyma adjacent to the blood vessels indicating brain tissue hemorrhage (arrows). Magnification 40x.
Deposition of immune complexes is a potent and rapid trigger for inducing vascular inflammation as it has the capacity to activate the complement pathway [5,39,53,61-63]. Indeed, inappropriate activation of the complement (i.e., in neurons and cerebral vasculature) is frequently observed in inflammatory neurodegenerative and neuroimmune diseases with underlying vascular dysfunction [53,61,63,64]. In particular, activation via the classical antibody-dependent complement pathway has long been recognized in immune complex-mediated diseases including vasculitis and SLE [37,39,65].

Vascular deposition of immune complexes and leukocyte recruitment to the vascular wall are themselves dependent on activation of the antibody-dependent complement pathway. In particular, abnormal expression of C1q by the vascular smooth muscle cells and vascular endothelium facilitates binding of T-lymphocytes and immune complexes to the vascular wall, resulting in vascular damage, ischemic lesions and brain edemas [53,62,66]. The demonstration of the crucial role of C1q activation in secondary brain edema due to compromised cerebral vasculature [53] is of particular relevance to Case 2 described in this report, where cerebral edema with bilateral uncal notching and early cerebellar tonsillar herniation was revealed on autopsy. Moreover, the autopsy of Case 2 also revealed ischemic changes affecting the Purkinje cells in the cerebellum which, according to the coroner’s report, were indicative of a terminal ischemic-hypoxic encephalopathy.

It has further been demonstrated that both vascular endothelial and smooth muscle cells express MHC II and thus by facilitating T-cell recruitment essentially operate as competent antigen-presenting cells [36,67,68]. In this regard it is important to note that lymphocyte trafficking through the CNS is normally limited and that lymphocyte adhesion to brain endothelium is very low (less than 5% compared with 15%-20% in other organs) [39]. Furthermore, although cerebral vascular endothelial cells are capable of expressing MHC class I and II molecules (which are crucial for antigen presentation to the T lymphocytes), this occurs less often than in endothelium of the systemic vasculature [39].

Thus, the prominent and consistent HPV-16L1 Ig-complex, CD3+ T-cell, MHC II, C5b-9 and C1q vascular staining patterns in the brain tissue specimens described here (Figures 3-6), strongly point to a vasculitic neuropathy, triggered by an aberrant hyper-active immune response, as a plausible explanation for the fatal outcomes post-HPV vaccination in these two cases. Specifically, all six markers prominently stain numerous immune cells adhering to the walls of cerebral blood vessels (Figures 1-6). In addition, C1q and HPV-16L1 prominently stain the vascular endothelium and smooth muscle cell layer (Figures 3 and 4). Collectively, these findings suggest the involvement of both immune complexes and cross-reactive HPV-16L1 antibodies (binding to vascular walls) in triggering vascular damage.

The dense and prominent immunoreactivity for the membrane attack complex (MAC) component C5b-9 in perivascular deposits (Figure 5) provides further corroborative evidence for a vasculopathy resulting from an abnormal activation of the classical antibody-dependent complement pathway. The formation of MAC is a terminal step in the classical pathway and its activation results in a lytic destruction of target cells [64,69]. In immune vasculopathies, C5b-9/MAC activity is associated with destruction of both perivascular tissues and cells within the vessel wall [37,39]. MAC is normally activated by immune triggers, including vaccinations [70,71]. Activation of MAC in the brain in the absence of an active brain infection due to an
Cerebral vasculitis-associated symptoms reported to VAERS post-HPV vaccination

<table>
<thead>
<tr>
<th>VAERS ID</th>
<th>Event Category</th>
<th>Adverse event description</th>
</tr>
</thead>
<tbody>
<tr>
<td>425345-1</td>
<td>Hospitalized Serous (cerebral vasculitis)</td>
<td>This case was received from the Health Authority on 02-JUN-2011. Reference number: ADR 21077360. This case was medically confirmed. A 17 year old female patient received the second dose of GARDASIL (manufacturer, lot and batch number not reported) i.m. on 30-DEC-2008. On an unspecified date in March 2009, approximately three months post vaccination, the patient developed headaches, weakness and lack of coordination. The patient had visual disturbances, nystagmus, weakness of the right arm (not reported) on 02-DEC-2008. The patient was eventually admitted to hospital in 2009 for long periods of time and was extensively investigated. The patient had a rare cerebral vasculitis of unknown cause. This was steroid responsive but the patient was unable to speak, walk or coordinate her movements. The patient was fed through a percutaneous endoscopic gastrostomy and was cared for 24 hours a day. At the time of reporting, the patient had not yet recovered (also reported as unknown). The reporter stated that whilst it clearly could not be proven that the patient's illness was directly linked to GARDASIL vaccine she had a serious disability. The patient's illness was rare and unexplained and her parents were convinced that it was caused by the vaccine. The events were considered serious due to hospitalization and disability. Other business partner number included: E2011-03389. No further information is available.</td>
</tr>
<tr>
<td>338235-1</td>
<td>Hospitalized Serous (cerebral vasculitis)</td>
<td>Information has been received from the foreign Health Authorities (Reference no.: PEI20080801988), concerning a 17 year old female patient, who on 18-SEP-2007, was vaccinated with the first dose of GARDASIL (toleration was not reported); on 20-NOV-2007, was vaccinated with the second dose of GARDASIL (toleration not reported) and on 11-MAR-2008, was vaccinated with third dose of GARDASIL, intramuscularly (Lot # and injection site not reported). Starting mid-Jun-2008, the patient developed cerebral vasculitis with acute psychiatric symptoms and seizures. Cerebral biopsy, performed on 26-Aug-2008, revealed astrocytic gliosis. Lab finding showed increased number of cells in cerebrospinal fluid (+CSF). Despite of therapies with cortisone, neuroleptics and antiepileptics, the patient only improved slowly from the severe organic brain syndrome with slowdown, cognitive disorders and emotional symptoms. The cause of cerebral vasculitis remained unexplained. The patient had not recovered.</td>
</tr>
<tr>
<td>396220-1</td>
<td>Life-threatening Serious (cerebral vasculitis)</td>
<td>Information has been received from a physician concerning her 15 year old daughter with no pertinent medical history or drug allergies who in September 2009, was vaccinated with a first dose of GARDASIL (lot # not reported). No concomitant medications were reported. It was reported that in September 2009, very soon after the injection of GARDASIL, the patient developed fatigue and nausea. When the patient was on leaving the physician's office she experienced syncope. The patient began having symptoms of high fever, headache and sore muscles. The patient was ilitergic for months after she received GARDASIL. In June 2010 the patient had chest pain and went to the Emergency Room where she was diagnosed with anxiety. The EKG showed arrhythmias. The patient became weak and had to quit the Track Team at school. In June 2010 the patient was tired and lethargic and began to forget things. Approximately three weeks ago, in approximately August 2010, the patient developed a tremor in her left hand and she was unable to speak. The patient developed chronic seizures with chorea. On 08-AUG-2010 the patient was admitted to hospital. On 15-AUG-2010 the patient developed a stroke and was non verbal. The patient was diagnosed with drug induced systemic lupus erythematosus with chorea. It was reported that there was central nervous system (CNS) involvement and the patient had vasculitis of the brain. The physician did not report what type of diagnostic testing the patient had received. The patient was being treated with ten different medications reported as follows: intravenous immune globulin (IVIG), antibiotics, anti hypertensive medications due to kidney involvement and high doses of IV prednisone. The physician stated that today, on 17-AUG-2010, the patient was to begin chemotherapy with IV cyclotoxin for a duration of 24 hours. Currently, the patient had her voice back and was able to move all of her extremities except the left arm. As of 16-AUG-2010 the patient had not recovered from drug induced systemic lupus erythematosus with chorea, anxiety and arrhythmias. The patient was presently in hospital. The reporter felt that drug induced systemic lupus erythematosus with chorea was related to therapy with GARDASIL. Drug induced systemic lupus erythematosus with chorea, anxiety and arrhythmias were considered to be disabling, another important medical event and immediately life-threatening. Additional information has been requested.</td>
</tr>
<tr>
<td>335698-1</td>
<td>Not Serious (cerebral vasculitis)</td>
<td>Information has been received from a physician, concerning a patient who was vaccinated with GARDASIL vaccine 0.5 ml, intramuscularly. The physician reported that after receiving the GARDASIL vaccine, about 2 months ago, the patient experienced cerebral vasculitis and seizures. The doctor was unsure if it was the first or second 0.5 ml dose. The doctor was not sure if the patient would be seeing a specialist about this AE. The outcome is unknown. The patient sought unspecified medical attention. Upon internal review, seizures were determined to be an other important medical event. Additional information has been requested.</td>
</tr>
<tr>
<td>436679-1</td>
<td>Hospitalized Serious (cerebral vasculitis)</td>
<td>Case received from the Health Authorities in a foreign country on 23-SEP-2011 under the reference number PP20110312. Case medically confirmed. A 14 year old female patient was vaccinated intramuscularly with the first and second doses of GARDASIL (batch number not reported) on 06-OCT-2010 and 15-DEC-2010. On 11-MAR-2011 she experienced vertigo and migraine during 15 to 18 hours. On 12-MAR-2011 brain MRI was performed to evaluate these symptoms (contrast was not administered). Brain MRI findings showed hyperintensities on white substance hypointensities, and medullar MRI was normal. On 25-JUL-2011 brain MRI showed persistence of the periventricular white substance hypointensities, stable. The patient was seen in neurology consultation which revealed numerous non inflammatory but vascular-type lesions, and no evident diagnosis. The third dose of GARDASIL was not administered. The patient's hospitalization dates were not provided. At the time of reporting the patient had not recovered. The Health Authorities assessed the causal relationship between the reported reactions and vaccination as doubtful (C2 S1 L1) according to the foreign method of assessment. Other business partner numbers included: E2011-05715.</td>
</tr>
</tbody>
</table>
Table 3: Sample of VAERS reports related to cases of vasculitis following HPV vaccination. The VAERS database [59] was searched using the following criteria: (1) Symptoms: vasculitis, vasculitis cerebral, vasculitis gastrointestinal, vasculitis necrotising; (2) Vaccine products: HPV2 (human papilloma virus bivalent), HPV4 (human papilloma virus types 6,11,16,18), HPV (human papilloma virus); (3) Event category: all events; (4) Territory: all locations; (5) Date vaccinated: 2006-2012 (HPV vaccine post-licensure period). A total of 40 events were reported to VAERS. Note however that the clinical presentation of vasculitis is highly variable with fluctuating signs and symptoms and it is thus probable that many cases remain unreported, undiagnosed and/or misdiagnosed.

<table>
<thead>
<tr>
<th>Case Number</th>
<th>Event Type</th>
<th>Vasculitis</th>
<th>Event Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>387696-1</td>
<td>Hospitalized</td>
<td>Serious</td>
<td>Vasculitis unspecified</td>
</tr>
<tr>
<td>400182-1</td>
<td>Not Serious</td>
<td>Not Serious</td>
<td>Vasculitis unspecified</td>
</tr>
<tr>
<td>374104-1</td>
<td>Not Serious</td>
<td>Not Serious</td>
<td>Generalized vasculitis</td>
</tr>
<tr>
<td>452887-1</td>
<td>Not Serious</td>
<td>Not Serious</td>
<td>Vasculitis unspecified</td>
</tr>
</tbody>
</table>

Invasion of foreign infectious agent (i.e., virus or bacteria) thus indicates an abnormal triggering of the immune response in which the immune attack is directed towards self-tissue. Such activation of the MAC in brain tissue specimens from Case 1 and Case 2 can therefore plausibly be explained by the receipt of the qHPV vaccine given that in both cases the autopsy examinations found no evidence of an alternative microbiological insult to explain the fatal outcomes. Moreover, prominent MAC immunoreactivity was also detected in cerebellar Purkinje cells and neurons in various areas of the brain, including the hippocampus. These results will be presented and discussed in greater detail in a separate publication.

In addition, we detected intense MMP-9 immunostaining in the vascular wall and the perivascular extracellular matrix (ECM) in brain tissue specimens from both cases. The vascular wall MMP-9 immunostaining pattern closely coincided with that of HPV-16L1 and C1q (Figures 3 and 4). This finding again corroborates our suggestion of an immune-triggered vasculopathy as an underlying cause for the fatal outcomes in both cases following qHPV vaccination. Indeed, MMP-9 is one of the key pro-angiogenic enzymes involved in progression of inflammatory infiltrates and ECM destruction in vasculitis [41-43]. Elevated MMP-9 expression is strongly associated with neuroimmune vasculopathies and diverse severe inflammatory nervous system pathologies, including ischemic nerve damage in SLE [41], autoimmune demyelinating syndromes, ischemia and stroke [72,73]. MMPs are also prominently elevated in neuropathic pain [41], autoimmune demyelinating syndromes, ischemia and stroke [72,73]. The obvious limitations of our study are that the tissues examined represent two individuals against which there were no control samples. For this reason, we could not obtain a quantitative measure of neuroinflammation [77,78] of the central and peripheral nervous systems. In particular, in bacterial meningitis, acute breakdown of the blood-brain barrier, intrathecal production of pro-inflammatory cytokines and accumulation of blood derived leukocytes in the cerebrospinal fluid lead to brain edema, cerebral vasculitis, and ultimately permanent neuronal injury. An overactive immune response of the host, rather than the bacterial pathogen per se, is thought to be the cause of neuronal injury, resulting in permanent neurological sequelae [43,79]. In this regard, it is important to re-emphasize the fact that vaccines, designed to hyper-stimulate the immune response (owing to the action of immune adjuvants), appear to carry an inherent risk for serious autoimmune disorders affecting the CNS [17,80-82]. Based on exhaustive investigations of post-vaccination induced autoimmune phenomena, Cohen, Carvalho and Shoenfeld have concluded that, "it seems that vaccines have a predilection to affect the nervous system" [1,83].

The obvious limitations of our study are that the tissues examined represent two individuals against which there were no control samples. For this reason, we could not obtain a quantitative measure of neuroinflammation. We aim in the future to further corroborate our findings by examining brain tissues from other cases of sudden and unexplained death following HPV vaccination, as well as control brain tissue from age-matched individuals who clearly died from non-vaccination related causes. Nonetheless, the marked resemblance in immunostaining patterns for all immunohistological markers in brain tissues specimens in the present two cases (i.e., compare Figures 1-4), as well as the similarity between their symptoms and those noted on VAERS reports related to post-HPV vaccination vasculopathies (some of which were medically ascertained cases; Tables 2 and 3), strongly support our present conclusions.

Conclusion

Any medicinal product (including vaccines) carries some risk of
Cerebral vasculitis-related symptoms reported to U.S. VAERS following HPV vaccination

<table>
<thead>
<tr>
<th>Symptom</th>
<th># reports</th>
<th>% total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Syncope</td>
<td>3127</td>
<td>14.2</td>
</tr>
<tr>
<td>Dizziness</td>
<td>3012</td>
<td>13.7</td>
</tr>
<tr>
<td>Headaches, migraines</td>
<td>2286</td>
<td>10.5</td>
</tr>
<tr>
<td>Nausea</td>
<td>2147</td>
<td>9.8</td>
</tr>
<tr>
<td>Fatigue</td>
<td>1100</td>
<td>5.0</td>
</tr>
<tr>
<td>Seizures</td>
<td>923</td>
<td>4.2</td>
</tr>
<tr>
<td>Tremors</td>
<td>531</td>
<td>2.4</td>
</tr>
<tr>
<td>Psychotic symptoms, anxiety</td>
<td>256</td>
<td>1.2</td>
</tr>
<tr>
<td>Cognitive disorder, amnesia, memory impairment</td>
<td>164</td>
<td>0.8</td>
</tr>
<tr>
<td>Vasculitis cerebral</td>
<td>5</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Table 4: Cerebral vasculitis and related symptoms reported to U.S. VAERS following vaccination with HPV vaccines Gardasil and Cervarix in the post-licensure period (June 2006-September 2012).

The VAERS database [59] was searched using the following criteria: (1) Symptoms: syncope, dizziness, headaches, migraines, nausea, fatigue, seizures, tremors, psychotc symptoms, anxiety, cognitive disorder, amnesia, memory impairment and vasculitis cerebral; (2) Vaccine products: HPV2 (human papilloma virus bivalent), HPV4 (human papilloma virus types 6,11,16,18), HPV (human papilloma virus); (3) Event category: all events; (4) Territory: United States; (5) Date vaccinated: 2006-2012.

adverse effects. However, unlike most medicinal products, vaccines are often administered to otherwise healthy individuals and this fact, according to the FDA, places significant emphasis on their safety [84]. The proper understanding of a true risk from vaccines is thus crucial for avoiding unnecessary ADRs [19]. In this regard, the fact that to date no solid tests or criteria have been established to determine whether adverse events are causally linked to vaccinations [6,16] should be a cause for concern.

In recent years it has become increasingly clear that vaccines may be a triggering factor for severe neurological manifestations of autoimmune etiology [1-13]. Some of these autoimmune phenomena may be explained by molecular mimicry whereby an antigen of a recombinant vaccine (i.e., HPV or hepatitis B vaccine) or of a live, attenuated virus (i.e., MMR) may resemble a host antigen and trigger autoimmune [5,6,83]. Owing to their structural resemblance, antibodies and auto-reactive T cells not only destroy the invading pathogen but also attack the host tissue. The data from the present study not only validate the molecular mimicry hypothesis of vaccine-induced autoimmune diseases, but also further expand on the proposed pathway, which most likely begins with the passage of vaccine-derived HPV-16L1 antibodies across the blood-brain barrier and the choroid plexus. Once in the CNS, some of these antibodies bind to neuronal host-antigen(s) due to molecular mimicry, such as vascular endothelial and/or smooth muscle cells. This further leads to a classical antibody-dependent complement pathway activation (Clq and MAC), resulting in destruction of blood vessel integrity, hemorrhages and ischemic tissue injury. The resultant blood-brain barrier breakdown allows further nondiscriminatory passage of immune cells and vaccine-derived immune complexes into the brain thus perpetuating the HPV vaccine triggered neurodestructive autoimmune process. The HPV-16L1 VLPs appear to contribute to this aberrant immune process by invading the CNS (most likely via a macrophage-dependent Trojan horse mechanism), and depositing on the walls of cerebral blood vessels.

Cerebral vasculitis is a serious disease which typically results in fatal outcomes when undiagnosed and left untreated [37]. The fact that many of the symptoms reported to VAERS following HPV vaccination are indicative of cerebral vasculitis, but are unrecognized as such (i.e., intense persistent migraines, syncope, seizures, tremors and tingling, myalgia, locomotor abnormalities, psychotic symptoms and cognitive deficits; Table 4) is thus a serious concern in light of the present findings. In particular, the fact that positive cerebral blood vessel wall immunoreactivity was observed with HPV-16L1 and not with HPV-18L1 antibody (Figure 1) suggests that HPV vaccines containing HPV-16L1 VLPs (including Gardasil and Cervarix) pose an inherent risk for triggering potentially fatal autoimmune vasculopathies and are therefore inherently unsafe for some individuals. Exactly which individuals might be more prone to developing a serious ADR following HPV vaccination is currently unknown. It is also unknown whether HPV vaccination can actually prevent cervical cancer cases since the current optimistic surrogate-marker based extrapolations have not been validated and appear to have arisen primarily from significant misinterpretation of existing data [25,85]. Because the HPV vaccination programme has global coverage, the long-term health of many women may be at risk against still unknown vaccine benefits. In conclusion, any case of sudden unexpected death occurring after HPV or other vaccinations should always undergo an exhaustive immunohistochemical study according to the methods presented in this report.

Acknowledgments

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References


