Diffuse and Localized SARS-CoV-2 Placentitis
Prevalence and Pathogenesis of an Uncommon Complication of COVID-19 Infection During Pregnancy

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Abstract: Coronavirus disease 2019 (COVID-19) infection in pregnancy has been associated with preterm delivery and preeclampsia. A less frequent and underrecognized complication is extensive placental infection which is associated with high rates of perinatal morbidity and mortality. The frequency, early pathogenesis, and range of lesions associated with this infection are poorly understood. We conducted a population-based study of placental pathology from all mothers with COVID-19 (n = 271) over an 18-month period delivering within our health system. The overall prevalence of diffuse severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) placentitis, as defined by typical histology and immunohistochemical (IHC) staining for SARS-CoV-2 spike protein, was 14.8/1000, but increased to 59/1000 in preterm births. We also identified 3 cases with isolated small foci of localized SARS-CoV-2 placentitis, characterized by focal perivillous fibrin and intervillositis, which illustrate the early pathogenesis and suggest that infection may be contained in some cases. Two other placental lesions were more common in mothers with COVID-19, high-grade maternal vascular malperfusion in preterm deliveries and high-grade chronic villitis at term (5/5 cases tested of the latter were negative by IHC for SARS-CoV-2). Additional investigation of diffuse and localized SARS-CoV-2 placentitis by IHC showed loss of BCL-2, C4d staining in surrounding villi, and an early neutrophil-predominant intervillosal infiltrate that later became dominated by monocyte-macrophages. We propose a model of focal infection of syncytiotrophoblast by virally infected maternal leukocytes leading to loss of BCL-2 and apoptosis. Infection is then either contained by surrounding fibrinoid (localized) or initiates waves of aponecrosis and immune activation that spread throughout the villous parenchyma (diffuse).

Key Words: placenta, COVID-19, pathology, SARS-CoV-2, intervillositis, apoptosis, necrosis, perivillous fibrin, syncytiotrophoblast

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Coronavirus disease 2019 (COVID-19), caused by the novel beta-coronavirus severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has led to a worldwide pandemic first appreciated in the United States in late January 2020.1,2 Early in the pandemic, concerns were raised regarding the possible adverse effects of viral infection on pregnant mothers and their offspring.3 These concerns centered around 3 potential issues: the known increased maternal susceptibility to severe respiratory infection associated with immune deviation during pregnancy, transplacental spread as seen with many other viruses, and severe neonatal morbidity due to the immaturity of the developing immune system at birth.4

Early results were somewhat reassuring demonstrating only a mildly increased maternal risk of severe infection, a relatively small increase in preterm deliveries, and only rare possible transplacental spread with the relatively mild and transient neonatal disease.5 However, shortly thereafter, reports of an uncommon severe diffuse placental infection associated with maternal SARS-CoV-2 infection, referred to as diffuse SARS-CoV-2 placentitis for the purpose of this study, began to appear in April 2020.6,7 Despite multiple case reports and small case series of this unique pattern of placental infection, many physicians and patients are unaware of this important placental lesion and its strong association with perinatal morbidity and mortality.

A partial explanation for the susceptibility of some placentas to SARS-CoV-2 infection is an expression of the viral receptor angiotensin-converting enzyme-2 (ACE2) and coreceptor TMSS on placental syncytiotrophoblast (ScT), the multinucleate trophoblastic cell lining the surface of chorionic villi at the fetomaternal interface.8 However, it is well-documented that ScT is exceptionally resistant to infection by most pathogens and the reason why a small subpopulation of mothers with COVID-19 develop severe infection while the remainder are apparently unaffected remains obscure.9,10

We now report the results of a prospective, population-based study of all placentas delivered to mothers...
with COVID-19 in our health system over an 18-month period beginning in May 2020. Our study had 3 primary goals: (1) to determine the population prevalence of diffuse SARS-CoV-2 placentitis, (2) to identify and characterize localized forms of placental infection, and (3) to better understand the pathogenesis of this unusual placental response. We approached this last goal through a review of all well-documented cases of diffuse SARS-CoV-2 placentitis and a comparative immunohistochemical (IHC) study of both localized and diffuse cases plus 2 non–COVID-related placental conditions with overlapping histologic features, chronic histiocytic intervillositis (CHI), and massive perivillous fibrin deposition (MPVF).

MATERIALS AND METHODS

Study Population

This was a prospective population-based study of all placentas delivered to mothers testing positive for SARS-CoV-2 by reverse transcription-polymerase chain reaction during pregnancy and delivering at 1 of the 5 hospitals providing obstetric services in the University Hospitals Health System between May 1, 2020, and October 31, 2021. Our health system covers a large geographic area in Northeast Ohio and provides health care to ~33% of its residents. A total of 271 placentas from mothers testing positive for SARS-CoV-2 were examined, representing 2.7% of the 10,163 deliveries occurring over this time period. The institutional review board with the following stipulations approved the study. Available clinical data was limited to that normally accompanying the placenta (admission history and physical and selected additional information regarding the delivery). We also had no access to the records of liveborn infants delivered to affected mothers. Maternal COVID-19 symptoms on admission were described in 111 patients, 98 mild-moderate and 13 severe. Six mothers were hospitalized for COVID-19-related pneumonia. Placentas were evaluated

![Image](https://www.ajsp.com/images/article/image.png)

**FIGURE 1.** Histologic features of diffuse SARS-CoV-2 placentitis (case 1) with confirmation by IHC staining for SARS-CoV-2 spike protein. A, Extensive villous agglutination with ScT necrosis, perivillous fibrin (left), and chronic intervillositis (right) (hematoxylin and eosin). B, Strong positive staining for SARS-CoV-2 capsid protein involving nearly all ScT within areas of villous agglutination (SARS-CoV-2 immunostain). C, Extensive perivillous fibrin and ScT necrosis; manifesting features intermediate between non–COVID-related villous infarction and MPVF (hematoxylin and eosin). D, ScT nuclei with features suggestive of apoptosis: nuclear condensation, membrane blebbing, and cell shrinkage (hematoxylin and eosin).
by 1 or more of the authors, all members of our perinatal service which examines over 4000 placentas per year. Criteria for the histologic diagnosis of diffuse SARS-CoV-2 placentitis were as established in prior publications and described in detail below. The first 100 cases were reviewed by 1 of the authors (R.W.R.) with prior experience through review of positive cases seen in consultation from other institutions. All cases of SARS-CoV-2 placentitis were confirmed by IHC for SARS-CoV-2 spike protein with appropriate positive (autopsy lung) and negative controls (see below), as specified in recent publications.11

Population-based Controls
A database of singleton placentas evaluated according to Amsterdam consensus criteria between January 2005 and December 2014 at our institution was utilized to compare the prevalence of placental lesions in the current cohort to placentas evaluated before the COVID-19 era.12 This database includes 8006 placentas with an overall submission rate of 40% to 50% of total deliveries occurring over this time period.

Definition and Histologic Grading of Other Placental Lesions
Other placental lesions observed in placentas from mothers with COVID-19 infection were separated into 4 categories based on the recent Amsterdam classification system, maternal vascular malperfusion (MVM), fetal vascular malperfusion (FVM), idiopathic chronic villitis (so-called villitis of unknown etiology [VUE]), and acute chorioamnionitis.13 These categories were further subdivided into low-grade and high-grade subgroups, as previously reported.12 In brief, MVM was defined by accelerated villous maturation with high-grade MVM having distal villous hypoplasia, multiple villous infarcts, or placental weight less than third percentile. FVM was defined by avascular villi with high-grade FVM having an average of either >15 avascular villi per slide or multiple fetal thrombi. VUE was defined by villous stromal chronic inflammation with high-grade VUE having >10 contiguous affected villi. Acute chorioamnionitis was defined by neutrophils in the chorionic portion of the placental membranes with high grade having neutrophils in the umbilical arterial wall and/or confluent inflammation of chorionic vessels.

IHC Studies
All immunostaining was performed by the IHC laboratory at University Hospitals Cleveland Medical Center. Briefly, unstained 4 μm sections on whole charged slides were prepared from paraffin blocks and baked for 30 minutes at 60°C. Primary antibodies were incubated at 37°C, and slides were counterstained onboard the automated instruments listed below. The OptiView Universal DAB Detection Kit (Roche Ventana) was used for all assays. Individual conditions for each antibody are as follows.
SARS-CoV-2 Spike Protein (1A9, 1/750 Dilution, 30 min; GeneTex)

IHC was performed following incubation with Bond Epitope Retrieval Solution II (Leica), an EDTA-based pH 9.0 solution at 100°C. Negative controls were placentas with CHI, and MPVF diagnosed before the COVID-19 era. Positive controls were lung specimens from COVID-19-infected patients obtained at autopsy and a commercial positive lung control (Bio SB, Santa Barbara, CA).

Other Antibodies

CD3 (clone LN10; Leica), CD68 (clone 514H12; Leica), and myeloperoxidase (MPO) (clone M9A5; Leica) were processed using a Bond III automated immunostainer following incubation with the Bond Epitope retrieval solution II. BCL-2 (clone 124; Roche Ventana), CD4 (clone SP91; Roche Ventana), and CD56 (clone 123C3; Roche Ventana) were processed with a BenchMark Ultra Automated Immunostainer (Roche Ventana) following incubation with C1 epitope retrieval solution, a Tris-based buffer with a slightly basic pH solution at 95°C.

Statistical Analysis

Comparison of categorical data regarding the prevalence of placental lesions in various subgroups was performed by χ² analysis with 2 df with significance level set at P-value <0.05.
SARS-CoV-2 placentitis are provided in Table 1. Timing of positive SARS-CoV-2 testing and clinical symptoms relative to the date of delivery correlated poorly with the histology of the localized lesions, a finding also observed in other reported cases of diffuse SARS-CoV-2 placentitis.5,14 Case 5 demonstrated what appears to be an early, exudative pattern characterized by focal acute intervillitis with perivillous fibrin (Fig. 2A). Positive staining for SARS-CoV-2 spike protein within this lesion was limited to a short strip of contiguous ScT nuclei and a few maternal leukocytes in the adjacent intervillous space (Fig. 2B). Higher magnification images show recently coagulated maternal blood and focal acute inflammation in areas contiguous to degenerating ScT (Figs. 2C, D). Case 6 demonstrated what appears to be an organizing pattern with 2 small well-circumscribed foci, each showing serpiginous streaks of positive staining for SARS-CoV-2 spike antigen emanating from a central nidus of necrotic ScT and surrounded by fibrinoid matrix and focal extravillous trophoblast (Figs. 2E, F). Case 7 demonstrated what appears to be a remote, involuted pattern, seen in 1 of 2 twin placentas delivered to an asymptomatic mother with a positive SARS-CoV-2 test 51 days before delivery. Histologically, a basal intervillous thrombus surrounded a small cluster of completely involuted villi that showed residual IHC staining for SARS-CoV-2 spike protein (Figs. 2G, H). Interestingly, this case also had foci of low-grade VUE distant from the intervillous thrombus that were negative for SARS-CoV-2 by IHC.

### Controls and Other Cases With Negative IHC Staining for SARS-CoV-2 Spike Protein

Twenty-nine of 271 study placentas (10.7%) were evaluated by IHC for expression of the SARS-CoV-2 spike protein, and 22 lacked positive staining. These IHC negative cases included twelve with perivillous thromboinflammatory lesions: 5 with lesions suspicious for localized infection as described above, 5 with high-grade VUE, and 1 case each from SARS-CoV-2-infected mothers with features of CHI and MPVF but lacking ScT necrosis. Ten placentas from COVID-19-infected mothers without thromboinflammatory lesions were also negative by IHC. 5 from mothers with COVID-19-related pneumonia, and 5 from mothers with mild-moderate symptoms. Control placentas with CHI and MPVF from the pre-COVID-19 era also, as expected, lacked positive staining.

### Relative Frequency of Other Patterns of Placental Injury by Grade and Gestational Age in Mothers With COVID-19 During Pregnancy

The relative frequencies of other important placental lesions stratified by histologic grade (see the Materials and methods section) were compared with gestational age-matched controls from our institutional database of over 8000 placentas evaluated in a 10-year study before the COVID-19 pandemic (Table 3).12 While prevalence figures for the comparison group may be artificially elevated due to submission bias, lesions more common in the cohort from mothers with COVID-19 are worthy of further consideration. The frequency of most of these other placental lesions was equivalent in mothers with COVID-19 and controls for both term or preterm births. However, 2 lesions, high-grade VUE in term placentas and high-grade MVM among preterm deliveries, were significantly increased among SARS-CoV-2-infected mothers.

### TABLE 3. Frequency of Other Major Patterns of Placental Injury by Gestational Age and Maternal SARS-CoV-2 Status

| Placental Lesions | ≥37 Weeks Gestational Age | <37 Weeks Gestational Age |
|-------------------|---------------------------|---------------------------|
|                   | SARS-CoV-2 Positive | Controls | SARS-CoV-2 Positive | Controls |
| N                 | 220                     | 5311       | 51                     | 2389       |
| MVM               |                          |            |                        |            |
| Grade 1           | 5.5*                    | 9.3        | 5.9                    | 13.6       |
| Grade 2           | 0.9                     | 4.9        | 11.8**                 | 4.6        |
| FVM               |                          |            |                        |            |
| Grade 1           | 6.4                     | 10.4       | 7.8                    | 6.4        |
| Grade 2           | 1.4                     | 1.0        | 0                      | 1.4        |
| VUE               |                          |            |                        |            |
| Grade 1           | 2.7                     | 4.1        | 0                      | 2.7        |
| Grade 2           | 5.9**                   | 3.2        | 3.9                    | 2.1        |
| Acute chorioamnionitis |                |            |                        |            |
| Grade 1           | 9.6                     | 14.8       | 7.8                    | 9.6        |
| Grade 2           | 3.6                     | 7.8        | 2.0                    | 12.2       |

*Column percentages.

**P < 0.050, χ².

### FIGURE 2. Histologic features of early/localized SARS-CoV-2 placentitis with confirmation by IHC staining for SARS-CoV-2 spike protein. A, Localized, exudative pattern (case 5): sharply circumscribed focus of recent perivillous villous fibrin with scattered inflammatory cells and focal villous agglutination (hematoxylin and eosin). B, Localized, exudative pattern (case 5): cytoplasmic staining for SARS-CoV-2 capsid protein of a short, well-circumscribed strip of viable ScT (a few IHC-positive maternal leukocytes are seen in the intervillous space, other similar positive cells not shown) (SARS-CoV-2 immunostain). C, Localized, exudative pattern (case 5): maternal leukocytes (neutrophils, activated macrophages, and an eosinophil) embedded in a recent intervillous blood clot with focal ScT necrosis in adjacent villi (hematoxylin and eosin). D, Localized, exudative pattern (case 5): focally intense perivillous neutrophilic exudate with early ScT degeneration (hematoxylin and eosin). E, Localized, organizing pattern (case 6): small perivillous fibrin plaque with focally necrotic ScT (hematoxylin and eosin). F, Localized, organizing pattern (case 6): serpiginous strands of central positivity for SARS-CoV-2 capsid protein (SARS-CoV-2 immunostain). G, Localized, involuted pattern (case 7): ghost-like outlines of involuted villi surrounded by fibrinoid matrix and peripheral extravillous trophoblast (right). Inflammatory cells and viable ScT are not identified. (hematoxylin and eosin). H, Localized, involuted pattern (case 7): persisting positive staining for SARS-CoV-2 capsid protein outlines the contours of involuted central vill; surrounding villi, fibrinoid, and extravillous trophoblast lack staining (SARS-CoV-2 immunostain).
IHC Analysis of the Necroinflammatory Response in Localized and Diffuse SARS-CoV-2 Placentitis and Non–COVID-19-related Conditions With Overlapping Features

We took advantage of the distinct histologic features in 2 of our localized cases and one of the diffuse cases to investigate the different types of host responses to SARS-CoV-2 infection and compare them to 2 non–COVID-19-related conditions with overlapping features, CHI and MPVF. Case 7, the remote, involuted lesion, lacked inflammation and viable ScT and was therefore not further evaluated. Diffuse SARS-CoV-2 placentitis (case 1) showed diffusely decreased BCL-2 staining in virally infected ScT at the center of areas of villous agglutination (Fig. 3A). A much smaller focus of decreased ScT BCL-2 expression was seen in case 5, the localized early, exudative lesion (Fig. 3B). Case 6, containing 2 small organizing, localized lesions, showed a complete absence of BCL-2 staining in affected villi (Fig. 3C).

Extensive ScT C4d deposition accompanied diffuse SARS-CoV-2 placentitis, but in contrast to BCL-2, staining was most prominent in viable villi surrounding the more necrotic areas (Fig. 3D). There was no C4d staining within the localized early, exudative lesion, but a few small foci of C4d positive ScT were seen in histologically normal villi a short distance away (Fig. 3E). The 2 small organizing lesions lacked any C4d staining (Fig. 3F). The inflammatory infiltrate in diffuse SARS-CoV-2 placentitis was composed primarily of CD68-positive macrophages (Fig. 3G) with a lesser component of MPO-positive neutrophils (Fig. 3H), while the localized early, exudative lesion (case 5) showed the inverse pattern, a predominance of MPO-positive neutrophils with lesser numbers of CD68 macrophages (Figs. 3I, J). The 2 small organizing, localized, lesions showed only scant degenerating MPO-positive neutrophils in the center and relatively rare CD68-positive macrophages at the periphery (Figs. 3K, L). Other pertinent results not illustrated were as follows. Scattered lesional CD3-positive T cells were observed in both localized and diffuse SARS-CoV-2 placentitis, while no lesional CD56-positive natural killer (NK) cells were seen in any of the cases tested. Abundant CD56-positive cells in the adjacent maternal decidua served as an internal positive control. Cases of CHI and MPVF not associated with SARS-CoV-2 infection were distinct. While positive for CD68-positive macrophages, CHI had no areas of decreased BCL-2 or positive C4d staining and few MPO-positive neutrophils. MPVF, on the other hand, had diffusely decreased BCL-2 staining, but only rare foci of C4d positivity and lacked CD68-positive monocyte-macrophages.

DISCUSSION

Despite its unique histopathologic features and many previous case reports and short case series, diffuse SARS-CoV-2 placentitis (also known as CHI with trophoblast necrosis and placental SARS-CoV-2 infection with extensive perivillous fibrin and chronic intervillitis) remains a largely unappreciated and serious complication of maternal COVID-19 infection during pregnancy. Our study provides a snapshot of the placental consequences of COVID-19 in women delivering in Northeast Ohio over the first 18 months of the pandemic. We examined placentas from all mothers with positive SARS-CoV-2 infection; ~3% of the over 10,000 deliveries occurring over this time period. Four of these placentas showed diffuse SARS-CoV-2 placentitis for an overall prevalence of 14.8/1000 infected mothers. Notably, the prevalence of diffuse placentalitis was higher in preterm deliveries, affecting ~59/1000 of all SARS-CoV-2-infected mothers delivering before 37 weeks. Worthy of mention, all 4 cases occurred during periods of high COVID-19 spread in our area (cases 1 to 3, April-May 2021, and case 4, late October 2021). Interestingly, 3 additional cases were identified in the 5 weeks following the closure of the study, a period of high SARS-CoV-2 transmission in Northeast Ohio with a predominance of the delta variant.

A number of characteristic clinical features of diffuse SARS-CoV-2 placentitis have emerged, as illustrated by the review of our 4 new cases and 56 other well-characterized cases in the literature. Placental disease shows little or no correlation with the timing and severity of maternal COVID symptoms. This could reflect the presence of persistent reservoirs of SARS-CoV-2 at other sites. Cases also show little correlation with the severity of COVID-19 symptoms. Many

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cases have arisen in asymptomatic mothers, and immunostaining of the placentas of 5 mothers with severe COVID-19-related pneumonia in our study revealed no cases of positivity for SARS-CoV-2. There is no strong association of placentalitis with severe underlying maternal diseases, although obesity and hypertension are relatively frequent. Despite the extensive disease, fetoplacental growth restriction is rare, suggesting that placentalitis probably develops relatively close to the time of delivery. The extent of disease, which can approach 90% in some cases, likely explains the strong association with adverse outcomes such as stillbirth, premature delivery, and neonatal distress. To what extent diffuse SARS-CoV-2 placentalitis accounts for the recently documented 4-fold overall increase in the rate of stillbirth in COVID mothers remains to be determined, but all 3 stillbirths occurring in mothers with COVID-19 in our hospital (1 in this report and 2 subsequent cases) had diffuse SARS-CoV-2 placentalitis. While there is no overall fetal sex predilection for the development of placentalitis, affected stillborns are more likely to be male. However, the significance of this observation is uncertain since male predominance for severe adverse outcomes is a common feature of many pregnancy complications. Importantly, despite extensive placental disease, transmission to the fetus appears to be uncommon, and liveborn infants rarely show clinical signs and symptoms of COVID-19 infection.

Early in the course of the study, we detected unusual focal placental lesions characterized by inter villous fibrin and increased neutrophils. These lesions bear a superficial resemblance to those seen in some cases of maternal bacterial sepsis at term and in severe chorioamnionitis in the second trimester. IHC staining for SARS-CoV-2 spike protein was positive in 3 of 8 such cases that were evaluated, defining a hitherto unrecognized, localized variant of SARS-CoV-2 placentalitis. We suggest that these 3 cases may represent different histologic stages of localized infection. Case 5, an early, exudative lesion with a single small focus of inter villous hemorrhage and small clusters of neutrophils, showed early ScT necrosis with one short strip of ScT testing positive for SARS-CoV-2 spike protein by IHC. Interestingly, this early strip-like staining pattern resembles that previously reported in an in vitro study of normal term villi infected with SARS-CoV-2 virus in organ culture and evaluated 72 hours post infection. A second, organizing pattern with 2 small lesions in close apposition to one another (case 6) showed agglutinated villi with focal central SARS-CoV-2 staining surrounded by fibrinoid matrix and extravillous trophoblast. The third, late involuted pattern was observed in our case 7, an asymptomatic woman who had tested positive for SARS-CoV-2 51 days before birth. Staining for SARS-CoV-2, in this case, was limited to a small focus of ghost-like villi surrounded by a large bland inter villous thrombus. These cases suggest that early lesions may sometimes be “walled-off” by a combination of fibrin-type (maternal coagulation-derived) and matrix-type (inter villous extravillous trophoblast-derived) fibrinoid deposition and later involute completely. Importantly, we cannot exclude that the 5 other, histologically similar localized lesions identified in our study may have had levels of viral antigen below the threshold of detection of our IHC assay or that there was the loss of antigenicity due to prolonged intervals before fixation.

A controversial topic in the previous literature is whether gestational COVID-19 infection during pregnancy is associated with any of the classic patterns of placental injury, as defined by the Amsterdam classification. Early reports found associations with maternal and FVM that were not observed in subsequent studies. We took advantage of our large institutional database to examine the relative frequency of these other lesions, stratified by grade and gestational age, in patients with and without COVID-19. While selection bias is unavoidable, it should be noted that our overall placental examination rate is high (40% to 50% of all deliveries). We found a significantly increased rate of high-grade MVM in the placentas of preterm infants and an increased rate of high-grade VUE in the placentas of term infants delivered from mothers with COVID-19. We favor prepregnancy risk factors or COVID-19-related increased susceptibility, rather than direct viral causation, as potential explanations for these findings. Of note, the 5 cases of high-grade VUE tested were all negative for IHC for SARS-CoV-2 spike protein.

Hematogenous infections of the placenta fall into several distinct groups. Best known are so-called TORCH infections, such as cytomegalovirus, which cause chronic villitis in the placenta, have a high rate of spread to the fetus, and are associated with congenital anomalies. A second group are infections, such as Zika virus, that cause only focal placental inflammation but also have a high rate of spread to the fetus with frequent malformations. The third group are antenatal infections, such as Parvovirus B19, where organisms cross the placenta and infect the fetus but do not cause placental inflammation. SARS-CoV-2 placentalitis falls into a fourth group of very uncommon placental infections (eg, Ebolavirus, Chlamydia psittici, and Coxiella burnetti), where inflammation and necrosis are most severe in the ScT lining the villi and in the adjacent maternal intervillous space. The unique feature of organisms in this fourth group is the ability to replicate within ScT, a cell type known to be unusually resistant to infection compared with both other placental cells and cells in the remainder of the body. Unusual aspects of SARS-CoV-2, not seen with any of the other hematogenous infections, are its low prevalence among infected mothers, its exclusive (not just predominant) localization to ScT, and its low rate of spread to the fetus.

Our IHC analysis revealed some interesting additional details regarding pathogenesis. First, we found that BCL-2 staining is decreased in ScT, even at apparently early stages of infection. This decrease may predispose to apoptosis, which has previously been documented in diffuse SARS-CoV-2 placentalitis by immunostaining for caspase-3. Interestingly, in vitro experiments of trophoblast in tissue culture have demonstrated a unique pattern where focal apoptosis can spread rapidly in waves to involve the entire cellular
This may reflect the fact that ScT is a syncytium without the cell-cell junctions that normally impede such spread in other cell types. Second, as previously reported, we found strong C4d staining in diffuse SARS-CoV-2 placentitis, but in our cases, this staining was largely restricted to the surrounding villi, rather than those directly infected by SARS-CoV-2 as indicated by positive IHC. Complement is activated in SARS-CoV-2 infections by both the classical (antigen-antibody complexes) and lectin (mannosylated SARS-CoV-2 S protein and MASP activation) pathways. Decreases in complement regulatory proteins in damaged ScT may further enhance local activation, and complement fragments such as C5a likely contribute to the brisk inflammatory response seen in diffuse cases. With regard to the inflammatory response, we found that the localized early, the exudative lesion showed both MPO-positive neutrophils and CD68-positive monocyte-macrophages with the former predominating while the later, organizing lesion contained only rare degenerating neutrophils and a few scattered surrounding monocyte-macrophages. Diffuse placentitis, on the other hand, was associated with abundant neutrophils and monocyte-macrophages, with the latter predominating. While a few scattered CD3-positive T cells were seen adjacent to all lesions, there were no CD56-positive NK cells. This is somewhat surprising for 3 reasons. First, NK cells are a prominent response in pulmonary COVID-19 infections. Second, abundant CD56-positive cells are frequent in the immediately responding cell in pulmonary COVID-19 infections. Second, what surprising for 3 reasons. First, NK cells are a prominent lesion, there were no CD56-positive NK cells. This is somewhat surprising for 3 reasons. First, NK cells are a prominent

We found that the localized early, the exudative lesion showed both MPO-positive neutrophils and CD68-positive monocyte-macrophages with the former predominating while the latter, organizing lesion contained only rare degenerating neutrophils and a few scattered surrounding monocyte-macrophages. Diffuse placentitis, on the other hand, was associated with abundant neutrophils and monocyte-macrophages, with the latter predominating. While a few scattered CD3-positive T cells were seen adjacent to all lesions, there were no CD56-positive NK cells. This is somewhat surprising for 3 reasons. First, NK cells are a prominent response in pulmonary COVID-19 infections. Second, abundant CD56-positive cells are frequent in the immediately adjacent decidua. Third, one study suggests that NK cells may play an important role in controlling hematogenous placental infection caused by *Listeria monocytogenes*.

Finally, CHI and MPVF unrelated to COVID-19 infection have some overlapping histologic features but have not been directly compared with SARS-CoV-2 placentitis in previous studies. We found their immunoprofile to be distinct with fewer neutrophils, normal BCL-2 and absent C4d staining in CHI and absent inflammation, and only focal C4d staining in MPVF. Of note, recent studies suggest that MPVF and CHI are phenotypes showing some overlap and that distinct clinical subsets may exist with different IHC profiles.

Based on the findings of this and previous studies, we propose the following working model for placentitis associated with maternal COVID-19 infection. First, placentitis is rare because of the known intrinsic ability of ScT to resist infection. We suggest that decreased resistance to infection involves loss of function in known protective pathways, such as high levels of interferon-delta, inflammasome activation, or C19 miRNA expression. Loss of function could be due to intrinsic genetic abnormalities, such as the interferon pathway mutations described in some adults, or to extrinsic factors such as high levels of circulating virus, upregulation of ACE2 viral receptor expression, or focal ScT hypoxia. An example of an intrinsic factor predisposing to the related placental lesion, MPVF, is fetal long-chain 3-hydroxyacyl-CoA dehydrogenase deficiency. An example of an extrinsic factor is the recently demonstrated colocalization of increased ACE2 receptor in areas of villous hypoxia as demonstrated by IHC for carbonic anhydrase IX. Once productively infected, ScT, despite its known high levels of endogenous autophagy, may be limited in its ability to kill the virus due to the known propensity of SARS-CoV-2 to persist and replicate in autophagocytic vesicles avoiding fusion with lysosomes. Infected ScT might then progress along 1 of 2 pathways: early complete apoptosis with surrounding fibrinoid deposition (localized SARS-CoV-2 placentitis) or uncontrolled waves of incomplete apoptosis (aponecrosis) resulting in a diffuse MPVF-like response. Interestingly, similar MPVF-type responses have recently been reported with other placental infections including as cytomegalovirus, coxsackievirus, and syphilis. While this diffuse MPVF-type pattern may partially protect the mother, and possibly the fetus, from high-level infection, it does so at the expense of extensive placental damage that can lead to adverse pregnancy outcomes. Persisting high levels of virus in ScT may lead to continuing generation of complement fragments, inflammasome-derived cytokines, and other chemotactic factors that act in a feed-forward fashion to elicit the monocyte-macrophage predominant inflammatory response that further aggravates placental dysfunction. These factors together could explain the unique destructive lesion known as diffuse SARS-CoV-2 placentitis.

Strengths of our study include its population-based design spanning the entire pandemic period up to November 2021, the uniform pathologic analysis by a single group of perinatal pathologists with extensive experience in evaluating a large number of placentas using criteria as defined by the Amsterdam consensus classification, and a large pre-COV-ID-19 placental database from the same patient population for comparisons. Weaknesses include selection bias in the comparison group since only 40% to 50% of all placentas were evaluated, the fact that only 11% of COVID-19 placentas were stained for SARS-CoV-2 spike protein by IHC, the intrinsic limitations of inferring causality and timing by analyzing pathologic specimens at any one instant in time, and the lack of neonatal follow-up.

From a practical perspective, we find that all placentas showing the 3 classical features, diffuse villous agglutination, ScT necrosis, and a histioocyte-predominant intervillous inclusions, are positive for SARS-CoV-2 by IHC, and hence a presumptive diagnosis can be provided to the clinician before, or even without, confirmation. In our experience, other patchy-diffuse lesions lacking 1 or more of the defining characteristics were never IHC positive. However, localized lesions containing SARS-CoV-2 spike protein by IHC exist and can show several distinct patterns. With experience, such lesions are easily identifiable by hematoxylin and eosin stain and can be further evaluated using either IHC or RNA in situ hybridization. However, given the limited availability of these tests and the absence of clinical data regarding clinical significance, the role of ancillary testing remains to be established.

In conclusion, we report an overall prevalence of 1% to 2% for diffuse SARS-CoV-2 placentitis in mothers with gestational COVID-19 infection and a 10-fold increase in preterm births. We define and characterize a novel localized form of SARS-CoV-2...
placentitis and provide some additional insights into the pathogenesis of both the localized and diffuse variants. We also report anecdotal evidence for increased prevalence during periods of COVID-19 surges in our area, especially with respect to the recent delta variant-related surge. Outstanding questions include further details regarding the molecular pathways leading to localized versus diffuse placental involvement and the frequency, pathogenesis, and long-term consequences of any fetal infections occurring in this setting.

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