Alagille syndrome and risk for hepatocellular carcinoma: Need for increased surveillance in adults with mild liver phenotypes

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Abstract
Alagille syndrome (ALGS) is a multisystem autosomal dominant developmental disorder caused predominantly by pathogenic variants in JAGGED1 (JAG1), and also by pathogenic variants in NOTCH2 in a much smaller number of individuals. Clinical presentation is highly variable and includes liver, heart, eye, skeleton, and facial abnormalities, with a subset of individuals also presenting with kidney, vascular, and central nervous system phenotypes. Hepatocellular carcinoma (HCC) is a rare complication of ALGS, though little is known about its incidence or etiology among affected individuals. Previous reports have identified HCC occurrence in both pediatric and adult cases of ALGS. We present a case report of HCC in a 58-year-old woman with a pathogenic JAG1 variant and no overt hepatic features of ALGS. Through a comprehensive literature review, we compile all reported pediatric and adult cases, and further highlight one previously reported case of HCC onset in an adult ALGS patient without any hepatic disease features, similar to our own described patient. Our case report and literature review suggest that ALGS-causing variants could confer risk for developing HCC regardless of phenotypic severity and highlight a need for a cancer screening protocol that would enable early detection and treatment in this at-risk population.

KEYWORDS
cholestasis, hepatocellular carcinoma, JAG1, liver disease, notch signaling

1 | INTRODUCTION

Alagille syndrome (ALGS, OMIM #118450), also known as syndromic bile duct paucity or arteriohepatic dysplasia, is a multisystem autosomal dominant developmental disorder with highly variable expression. This syndrome was first reported by Alagille et al. in 1969 with an emphasis on the hepatic manifestations, and has since been identified as the most common form of genetic cholestatic liver disease (Alagille et al., 1987; Alagille, Habib, & Thomassin, 1969). ALGS is also characterized by abnormalities of the heart, eye, and skeleton, including a characteristic facial appearance, as well as less common features involving the kidneys, vasculature, and central nervous system (Emerick et al., 1999; Emerick et al., 2005). The disease has been recognized as a dysfunction of the Notch signaling pathway, an integral developmental process involved in cell fate determination (Huang, Li, Zheng, & Wei, 2019; Morell, Fiorotto, Fabris, & Strazzabosco, 2013). The majority of ALGS-associated pathogenic variants (94.3%) have been identified in the JAGGED1 (JAG1) gene, which encodes a Notch
ligand, with a smaller percentage of pathogenic variants (2.5%) identified in the NOTCH2 gene, which encodes a Notch receptor (Gilbert et al., 2019; Kamath et al., 2012; Li et al., 1997; McDaniell et al., 2006; Oda et al., 1997; Saleh, Kamath, & Chitayat, 2016). A small number (3.2%) of individuals meet the clinical criteria for ALGS, but do not have a pathogenic variant detected in either disease-causing gene (Gilbert et al., 2019).

To date, 694 pathogenic variants in JAG1 and 19 pathogenic variants in NOTCH2 have been described in individuals with clinical features of ALGS (Gilbert et al., 2019). JAG1 variants are most typically protein-truncating, and haploinsufficiency is believed to be the disease-causing mechanism (Oda et al., 1997; Saleh et al., 2016; Spinnert et al., 2001). The mechanism by which NOTCH2 variants cause ALGS is not clear, as only 19 have been described, although limited functional analysis has demonstrated reduced Notch signaling for a handful of described variants (Gilbert et al., 2019; Kamath et al., 2012).

No studies have shown a definitive correlation between type of mutation and ALGS phenotype or severity (Crosnier et al., 1999; McDaniell et al., 2006; Mitchell, Gilbert, & Loomes, 2018; Spinnert et al., 2001).

Isolated case reports point to an association of hepatocellular carcinoma (HCC) among individuals with ALGS (Békássy, Garwicz, Wiebe, Hägerstrand, & Jensen, 1992; Bhadri et al., 2005; Chiaretti, Zampino, Botto, & Polidori, 1992; Kim et al., 2005; Pérez Becerra et al., 1991; Rabinovitz, Imperial, Schade, & Van Thiel, 1992; Syed, Khalili, & Guindi, 2008; Tsai et al., 2010; Valampampiril, Shannmugam, Vij, Reddy, & Rela, 2020; Wegmann, Evison, Schaub, Kist, & Vest, 1996). HCC has been documented in children and adults with ALGS phenotypes ranging from mild to severe and with differing degrees of liver involvement. The occurrence of HCC in multiple individuals with subtle clinical features (and some with little to no apparent liver involvement) has caused some to suggest that ALGS-causing variants could contribute to the development of HCC by interfering with Notch signaling. Abnormalities in Notch signaling have previously been implicated in HCC and intrahepatic cholangiocarcinoma, and mutations in JAG1 and NOTCH2 have been identified in a variety of human tumor tissues (Alagille et al., 1987; Alagille, Odievre, Gautier, & Dommergues, 1975; Emerick et al., 1999; Saleh et al., 2016). In all cases, HCC was confirmed by imaging and/or pathology. In most cases, malignancy was also reflected by elevated serum α-fetoprotein levels.

A comprehensive list of JAG1 and NOTCH2 variants previously identified in human tumor tissues, including HCC, was curated using the Catalogue of Somatic Mutations in Cancer (COSMIC) database (cancer.sanger.ac.uk; Tate et al., 2019). Allele frequency was obtained for each variant using the Genome Aggregation Database (gnomAD; https://gnomad.broadinstitute.org; Karczewski et al., 2020).

All included individuals were confirmed to have intrahepatic bile duct paucity and three of the five clinical criteria established by Alagille et al., an affected first-degree family member and two of the clinical criteria, and/or a confirmed JAG1 or NOTCH2 mutation (Alagille et al., 1987; Alagille, Odievre, Gautier, & Dommergues, 1975; Emerick et al., 1999; Saleh et al., 2016). In all cases, HCC was confirmed by imaging and/or pathology. In most cases, malignancy was also reflected by elevated serum α-fetoprotein levels.

Ethical considerations

This study was approved by the Institutional Review Board at the Children’s Hospital of Philadelphia, and written informed consent was obtained from parents or subjects 18 years or older in accordance with the Declaration of Helsinki.

RESULTS

Case report

A 58-year-old woman with ALGS presented with a persistent productive cough, reflux, and weight loss and was referred for complete abdominal ultrasound as well as frontal and lateral radiographs of the chest. She was diagnosed with ALGS around age 35 by molecular confirmation of a pathogenic JAG1 variant, identified in genomic DNA from whole blood, after the same variant (c.693_694del; p. Arg231Serfs*10) was identified in her two sons (Li et al., 1997, see...
Both sons had classic features of ALGS, including severe cholestasis and progressive liver disease requiring transplantation, whereas she presented with mild clinical features and had no history of hepatic complications. Despite her mild presentation, we do not have evidence to suggest that the proband was mosaic for the JAG1 variant (Li et al., 1997). Phenotypic features of the proband and her sons are summarized in Table 1.

An abdominal ultrasound revealed a large heterogeneous hypervascular mass in the right lobe of the liver measuring $10.8 \times 12.9 \times 11.8$ cm as well as mild bilateral renal cortical atrophy. The mass was initially presumed to be a cavernous hemangioma, with HCC also in the differential diagnosis. Subsequent multiplanar magnetic resonance imaging (MRI) with contrast performed 4 months later revealed that the mass had grown to measurements of $14.3 \times 10.9 \times 14.2$ cm, occupying most of the right hepatic lobe, suspicious for hepatocellular neoplasm (Figure 1). The proband had no history of known risk factors for developing HCC, such as obesity, alcoholism, Hepatitis B infection (HBV), Hepatitis C infection (HCV), or non-alcoholic fatty liver disease (NAFLD; Desai, Sandhu, Lai, & Sandhu, 2019).

At the time of the MRI, the proband presented with elevated gamma-glutamyltransferase as well as abnormal levels of albumin, direct bilirubin, alkaline phosphatase, aspartate aminotransferase (AST), and alanine aminotransferase (ALT), summarized in Table 2. Serum levels of tumor marker α-fetoprotein were within normal range. The MRI was consistent with HCC and she was referred to Hepatology and Gastroenterology specialists for management of
related complications. The size of the tumor at initial diagnosis precluded consideration of liver transplantation (LT) as a therapeutic option. The proband declined chemotherapy, ablation, or other therapeutic interventions, and no biopsy or tumor resection was performed. Somatic cancer testing was therefore unavailable.

She presented with abdominal pain 6 months later, at which time a second MRI with contrast revealed that the mass had grown to measurements of $15.6 \times 11.3 \times 16.6$ cm, exerting a stable mass effect on the right kidney. A repeat hepatic function panel obtained at this time revealed abnormal albumin, bilirubin, alkaline phosphatase, AST, and ALT (Table 2). Of note, serum \( \alpha \)-fetoprotein levels remained within normal range throughout the course of illness. Alkaline phosphatase, AST, and ALT levels continued to increase over time (Table 2). Five months after the second MRI, the proband presented to the emergency department with acute kidney injury, hyperkalemia, and ascites. She was transitioned to home palliative care the following day and died 5 months later at the age of 59, 15 months following diagnosis.

### 3.2 Literature review

We identified 28 publications describing 21 cases of HCC in children with AGLS, and 13 in adults (Tables 3 and 4). All pediatric patients with a described liver phenotype had paucity of intrahepatic bile ducts and cholestasis prior to developing HCC. The majority of adult patients with a described liver phenotype also had bile duct paucity and cholestasis, with the exception of one patient who, like our proband, had an attenuated form of ALGS with no overt liver involvement (Keeffe et al., 1993). Most pediatric cases arose on a background of cirrhotic liver (\( n = 16 \), of 18 cases that described presence or absence of cirrhosis). In contrast, only a few adult cases arose on a background of cirrhosis (\( n = 4 \), of nine cases that described this information). The presence and severity of other syndromic features, including characteristic facies and eye, heart, or skeletal abnormalities, was highly variable in both children and adults.

ALGS was identified prior to the diagnosis of HCC in the majority of cases, with the exception of one patient who died of HCC, where ALGS was diagnosed at postmortem evaluation (Wegmann et al., 1996).

Many of the reports pre-dated the discovery of \( JAG1 \) as the primary disease gene in 1997, and genetic data, obtained from screening genomic DNA extracted from whole blood, were available for only five patients, including the proband. Three had confirmed pathogenic \( JAG1 \) variants, including: an unspecified de novo variant in a protein-coding region (Patient 12, Table 3), the previously described, known pathogenic, missense variant c.551 G > A; p.Arg184His (Patient TABLE 2 Summary of abnormal laboratory test results in the proband

| Test                      | Normal range | At initial MRI | Time since initial MRI |
|---------------------------|--------------|----------------|------------------------|
|                           |              | 2 months       | 6 months               | 15 months              |
| Hepatic function panel    |              |                |                        |                        |
| Albumin (g/dl)            | 3.6–5.1      | 2.9            | 3.3                    | 2.9                    | 2.7                    |
| Total bilirubin (mg/dl)   | 0.2–1.2      | 0.9            | 0.8                    | 1.3                    | 1.5                    |
| Direct bilirubin (mg/dl)  | ≤0.2         | 0.3            | 0.2                    | 0.6                    | 0.8                    |
| Alkaline phosphatase (U/L)| 33–130       | 237            | 286                    | 869                    | 1,256                  |
| AST (U/L)                 | 10–35        | 309            | 108                    | 119                    | 265                    |
| ALT (U/L)                 | 6–29         | 142            | 38                     | 60                     | 102                    |
| GGT (U/L)                 | 3–70         | 250            | –                      | –                      | –                      |
| Tumor marker              |              |                |                        |                        |                        |
| AFP (ng/ml)               | <6.0         | 3.1            | 3.7                    | 3.5                    | –                      |

Abbreviations: AFP, alpha-fetoprotein; ALT, alanine aminotransferase; AST, aspartate aminotransferase; GGT, gamma-glutamyltransferase.

TABLE 5 Summary of \( JAG1 \) variants identified in human HCC samples

| Sample | Variant | Amino acid change | Reported in ALGS? | gnomAD frequency $^a$ |
|--------|---------|-------------------|-------------------|-----------------------|
| 1      | c.48C > A | p.(=)            | No                | $2.31 \times 10^{-5}$ |
| 2      | c.82-85C > G | p.?             | No                | Absent                |
| 3      | c.429 T > C | p.(=)            | No                | Absent                |
| 4      | c.886 + 94A > C | p.?          | No                | Absent                |
| 5      | c.2889del | p.Phe963Leufs*7 | No $^b$           | Absent                |
| 6      | c.3647A > G | p.Tyr1216Cys    | No                | Absent                |

Note: http://cancer.sanger.ac.uk.

$^a$http://gnomad.broadinstitute.org.

$^b$Although not reported in ALGS, this frameshift would be expected to cause ALGS. It is possible that this patient had undiagnosed ALGS.
| Patient | Age (years) | AFP (ng/ml) | Cirrhosis (Y/N) | Bile duct paucity (Y/N) | Cholestasis (Y/N) | Other features | Family history | Genetic findings | Treatment/outcome | Reference |
|---------|-------------|-------------|----------------|-------------------------|------------------|---------------|----------------|----------------|------------------|------------|-----------|
| 1       | 3.5         | 31,800      | Y              | Y                       | Y                | Facies, pulmonic stenosis, pruritus, jaundice, xanthomas, growth delay | Mother and maternal grandmother with ALGS. Mother had pulmonic stenosis and no neonatal cholestasis. | NR             | Tumor invasion precluded LT; patient died | Kaufman et al., 1987; Ong, Williams, Anderson, & Kaplan, 1986 |
| 2       | 16          | NR          | Y              | Y                       | Y                | Facies, pulmonic stenosis, pruritus, jaundice, delayed puberty, history of giant cell hepatitis | Brother and sister with ALGS and HCC. Second sister with ALGS and suspected HCC. | NR             | Died of variceal hemorrhage | Rabinovitz et al., 1989 |
| 3       | 2           | NR          | Y              | NR                      | NR               | Facies, systolic murmur, jaundice | Brother and sister with ALGS and HCC. Second sister with ALGS and suspected HCC. | NR             | Metastasis precluded LT; patient died | Rabinovitz et al., 1989 |
| 4       | 4           | 927.3       | Y              | Y                       | Y                | Facies, vertebral abnormalities, posterior embryotoxon, pulmonic stenosis, pruritus, jaundice | NR             | NR             | Died waiting for LT | Békássy et al., 1992 |
| 5       | 7           | NR          | NR             | Y                       | Y                | Facies, xanthomas | NR             | NR             | NR; patient died from rupture of HCC | Castañeda et al., 1992 |
| 6       | 17          | 400         | N              | Y                       | Y                | Facies, posterior embryotoxon, pulmonic stenosis, growth delay | NR             | NR             | Chemotherapy; patient died | Chiaretti et al., 1992 |
| 7       | 1.4         | 2,890       | Y              | NR                      | NR               | Facies, posterior embryotoxon, pulmonic stenosis, jaundice | No family history | NR             | LT; stable 20 months post-op | Kim et al., 2005 |
| 8       | 4           | 414         | Y              | Y                       | NR               | Facies, vertebral abnormalities, pulmonic stenosis, jaundice | No family history | NR             | No treatment; patient died | Kim et al., 2005 |
| Patient | Age (years) | AFP (ng/ml) | Cirrhosis (Y/N) | Bile duct paucity (Y/N) | Cholestasis (Y/N) | Other features | Family history | Genetic findings | Treatment/outcome | Reference |
|---------|-------------|-------------|-----------------|-------------------------|-------------------|----------------|---------------|----------------|----------------|-------------|
| 9       | 7           | 344.43      | Y               | Y                       | NR                | Facies, vertebral abnormalities, growth delay, corneal opacity, pruritus, jaundice | Younger brother with growth delay but no confirmed ALGS | NR             | No treatment; patient died | Kim et al., 2005 |
| 10      | 4           | 264,200     | N               | NR                      | Y                 | Facies, pulmonic stenosis, pruritus, jaundice | No family history | NR             | No treatment; patient died | Bhadri et al., 2005 |
| 11      | 4           | 8,155       | Y               | NR                      | Y                 | Facies, vertebral abnormalities, pulmonic stenosis, pruritus | Father and father’s sister with ALGS. Father’s sister died of associated congenital heart disease. | No JAG1 pathogenic variant | LT; died 3 months post-op of Escherichia coli septicemia | Bhadri et al., 2005 |
| 12      | 3           | NR          | Y               | Y                       | NR                | Facies, vertebral abnormalities, pulmonic stenosis, jaundice | No family history | de novo pathogenic JAG1 variant in a protein-coding region | LT; outcome NR | Wetli, Gralla, Schibli, & Stranzinger, 2010 |
| 13      | 15          | 1           | Y               | NR                      | NR                | NR             | NR            | Transarterial chemoembolization | Pham, Gallo, Concepcion, Esquivel, & Bonham, 2015 |
| 14      | 8           | 1,650       | Y               | NR                      | NR                | NR             | NR            | LT; alive | Geramizadeh et al., 2017 |
| 15      | 15          | 2           | Y               | NR                      | NR                | NR             | NR            | Transarterial chemoembolization followed by LT; alive | Weiss et al., 2018 |
| 16      | 13          | 2.1         | Y               | NR                      | NR                | NR             | NR            | LT; alive | D’Souza et al., 2018 |
| 17      | 1.5         | 3.1         | Y               | Y                       | Y                 | Facies, vertebral abnormalities, pulmonic stenosis, xanthomas, decompensated liver disease | No family history | JAG1 c.551G > A (p.Arg184His) | LT; alive | Valamparampil et al., 2019; Valamparampil et al., 2020 |
| 18      | 1.7         | 2           | Y               | Y                       | NR                | Aortic stenosis, xanthomas, decompensated liver disease | No family history | NR             | LT; alive | Valamparampil et al., 2019; Valamparampil et al., 2020 |
and the frameshift variant c.693_694del; p. Arg231Serfs*10 described in the proband (Table 1). One patient had a suspected pathogenic NOTCH2 variant, c.5830G > A (p.Gly1944Ser; Patient 19, Table 3). Genetic testing revealed no pathogenic variant in one patient, despite having a family history of ALGS (Patient 11, Table 3). Four of these five patients had severe liver disease, with the proband being the exception. Of the patients with confirmed JAG1 or NOTCH2 variants, only the proband had a family history of ALGS (Table 1). Somatic cancer testing of tumor samples was not available in any of the case reports or for our proband.

When the additional reports that do not contain genetic data are added to this group, a total of seven patients (20%) had a family history of ALGS, including the proband. Three of these were siblings with severe ALGS who all developed HCC (Patients 2 and 3, Table 3; Patient 24, Table 4). Eight patients (23%) had no family history of ALGS and this information was unavailable for the remaining 20 patients (57%; Tables 3 and 4).

### 3.3 Somatic variant review

A search of the Catalogue of Somatic Mutations in Cancer (COSMIC) database (cancer.sanger.ac.uk) yielded 719 human tumor samples (out of 38,357 sequenced) with JAG1 variants (1.9%) and 2,457 human tumor samples (out of 59,346 sequenced) with NOTCH2 variants (4.1%). JAG1 variants have been described in 28 unique tissue types, with variants in the liver, breast, large intestine, endometrium, and skin occurring at the highest frequency (Tate et al., 2019). Of those variants, 44.9% were missense, 3.9% were nonsense, and 3.4% were frameshift. Other types of JAG1 variants described in cancer are summarized in Table S1. Of the 20,411 liver tumor samples included in COSMIC, 3,289 were obtained from HCC. Six HCC samples had JAG1 variants and 24 had NOTCH2 variants (Tables 5 and 6; Tate et al., 2019). No JAG1 or NOTCH2 variants described in human HCC samples have also been identified in individuals with ALGS. However, the JAG1 frameshift p.Phe963Leufs*7 reported in Table 5 would be expected to cause ALGS and it is possible that this patient had undiagnosed ALGS.

### 4 DISCUSSION

We describe an individual with mild ALGS who presented with HCC and no overt ALGS liver phenotype. In addition, we highlight a similar case report of a second adult with mild ALGS presenting with HCC who also did not have an overt ALGS liver phenotype (Keeffe et al., 1993). Though their incidence is unknown, hepatic lesions are believed to be rare complications of ALGS (Rapp, Bellah, Maya, Pawel, & Anupindi, 2017). Lesions can manifest as regenerative nodules, a form of compensatory hyperplasia, or as HCC. Regenerative nodules are relatively benign, homogenous masses that do not usually necessitate clinical intervention (Alhammad, Kamath, Chami, Ng, & Chavhan, 2016; Bhadri et al., 2005; Ennaifer et al., 2016; Rapp

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### Table 3 (Continued)

| Patient | Age (years) | AFP (ng/ml) | Cirrhosis (Y/N) | Cholestasis (Y/N) | Bleed duct paucity (Y/N) | Vertebra anomalies (Y/N) | Pulmonary stenosis (Y/N) | Depressed liver (Y/N) | Other features | Treatment/outcome | Reference |
|---------|-------------|-------------|-----------------|------------------|-------------------------|-------------------------|------------------------|---------------------|--------------|----------------|-----------|
| 19      | 2           | 1           | Y               | Y                | Y                       | NR                      | NR                     | NR                  | Vertebral abnormalities, pulmonic stenosis, decompensated liver disease | LT; alive | D’Souza et al., 2020 |
| 20      | 9           | NR          | Y               | NR               | NR                      | NR                      | NR                     | NR                  | Neoadjuvant chemo followed by LT; died from dialysis complication | Neoadjuvant chemo followed by LT; died from relapse in lung and abdomen | D’Souza et al., 2020 |
| 21      | 3           | NR          | Y               | NR               | NR                      | NR                      | NR                     | NR                  | Neoadjuvant chemo | Neoadjuvant chemo | D’Souza et al., 2020 |

Abbreviations: AFP, α-fetoprotein; ALGS, Alagille syndrome; HCC, hepatocellular carcinoma; LT, liver transplantation; NR, not reported.
### TABLE 4  Summary of HCC cases reported in adult patients with ALGS

| Patient | Age (year) |AFP (ng/ml) | Cirrhosis (Y/N) | Bile duct paucity (Y/N) | Cholestasis (Y/N) | Other features | Family history | Genetic findings | Treatment/outcome | Reference |
|---------|------------|------------|-----------------|-------------------------|-------------------|----------------|----------------|----------------|-------------------|------------|
| 22      | 36         | 275        | N               | Y                       | Y                 | Pruritis, jaundice | NR             | NR              | Chemotherapy; outcome NR | Adams, 1986 |
| 23      | 31         | NR         | NR              | Y                       | NR                | Facies, vertebral abnormalities, posterior embryotoxon, pulmonic stenosis | NR             | NR              | Metastasis precluded LT; patient died | duCret, Cefalu, Alford, Drake, & Boudreau, 1988 |
| 24      | 25         | 207,000    | N               | Y                       | Y                 | Facies, posterior embryotoxon, pulmonic stenosis, jaundice, growth delay, skeletal abnormalities | Two brothers with ALGS and HCC. Sister with ALGS and suspected HCC. | NR             | Died of variceal hemorrhage | Rabinovitz et al., 1989 |
| 25      | 48         | 2,800      | N               | Y                       | NR                | Facies, previous HCC (resected), jaundice (spontaneous resolution in adolescence) | NR             | NR             | NR; patient died | Bail et al., 1990 |
| 26      | 43         | NR         | NR              | Y                       | NR                | NR             | Son with ALGS | NR             | NR               | Perez Becerra et al., 1991 |
| 27      | 31         | NR         | N               | Y                       | NR                | Facies, posterior embryotoxon, pulmonic stenosis, developmental delay, growth delay, pruritus | NR             | NR             | NR; patient died | Schwarzenberg et al., 1992 |
| 28      | 46         | 901        | NR              | N                       | N                 | Facies         | NR             | LT; outcome NR | Keefte et al., 1993 |
| 29      | 31         | NR         | NR              | NR                      | NR                | NR             | NR; ALGS not diagnosed until postmortem evaluation of medical history | NR             | NR             | NR; patient died | Wegmann et al., 1996 |
| 30      | 42         | NR         | Y               | Y                       | NR                | Pruritus       | NR             | NR             | NR               | Syed et al., 2008 |
| 31      | 29         | 2,500      | N               | Y                       | NR                | Facies, pulmonic stenosis | NR             | NR             | Successful resection | Tsai et al., 2010 |
| 32      | 25         | NR         | Y               | NR                      | NR                | NR             | LT; patient died of sepsis | Vinayak et al., 2017 |
| 33      | 41         | 3,7        | Y               | Y                       | NR                | NR             | Transarterial chemoembolization followed by LT due to recurrence; alive | Galvez et al., 2020 |
| 34      | 38         | 187        | Y               | NR                      | NR                | NR             | NR             | Right hepatectomy; patient died of infectious complications | Schoen, Porta, & Horvat, 2021 |

Abbreviations: AFP, α-fetoprotein; ALGS, Alagille syndrome; HCC, hepatocellular carcinoma; LT, liver transplantation; NR, not reported.
et al., 2017; Syed et al., 2008; Wettl et al., 2010). On the other end of the spectrum, HCCs are malignant, characterized by one or more heterogeneous tumor(s) prone to necrosis, hemorrhage, and metastasis (Kaufman et al., 1987; Kim et al., 2005; Syed et al., 2008). Malignancy is often, but not always (as in the case of our proband), reflected in elevated serum α-fetoprotein as well. HCCs are largely resistant to chemotherapy; however, extensive invasion of healthy liver tissue often precludes resection or transplantation and the probability of recurrence is high (Bail et al., 1990; Bhadri et al., 2005; Desai et al., 2019; Kaufman et al., 1987; Viatour et al., 2011). Screening and early intervention are therefore essential for a favorable prognosis.

It was initially hypothesized that HCC developed as a result of cholestasis associated with ALGS (Békássy et al., 1992; Chiaretti et al., 1992). However, there is no evidence that a seemingly milder ALGS phenotype correlates with a reduced risk for developing HCC. Our observation of HCC in two mildly-affected patients (i.e., without overt liver involvement) points to a possible disease association, with the argument that ALGS-causing germline variants could predispose affected individuals to HCC by interfering with Notch signaling (Bhadri et al., 2005; Tsai et al., 2010). Notch signaling is important in liver disease, where it has been shown to signal hepatic progenitor cell differentiation in the setting of cirrhosis with increased necroinflammation and fibrosis (Huang et al., 2019; Morell et al., 2013). Extensive upregulation of this repair pathway can result in malignant transformation of hepatic progenitor cells into cancer stem cells (Huang et al., 2019; Morell et al., 2013). Additionally, JAG1 is the predominant ligand stimulating Notch activation in injured livers (Fabris et al., 2007; Geisler & Strazzabosco, 2015; Morell et al., 2013). In JAG1 pathogenic variant-positive ALGS, the absence of functional protein results in a buildup of hepatic progenitor cells with an intermediate hepatobiliary phenotype incapable of differentiating into biliary cells (Fabris et al., 2007). Inability to repair damage could facilitate the progression of preexisting liver disease and subsequent development of HCC.

Notch signaling dysfunction is already well-described in cancer, with perturbations in Notch signaling found in solid liver, pancreatic, breast, and ovarian tumors, as well as in melanoma and glioblastoma (Lu et al., 2016; Tate et al., 2019). However, whether Notch signaling plays an oncogenic or tumor-suppressing role depends on the tissue type, surrounding microenvironment, and cooperation with other

**TABLE 6** Summary of NOTCH2 variants identified in human HCC samples

| Sample | Variant | Amino acid change | Reported in ALGS? | gnomAD frequencya |
|--------|---------|------------------|------------------|------------------|
| 1      | c.61G > A | p.Ala21Thr      | No               | 0.115            |
| 2      | c.155 + 5264A > G | p.?              | No               | Absent           |
| 3      | c.155 + 10578C > T | p.?              | No               | Absent           |
| 4      | c.344G > A | p.Cys115Tyr     | No               | Absent           |
| 5      | c.415 + 14C > T | p.?              | No               | Absent           |
| 6      | c.416-2,629 T > C | p.?              | No               | Absent           |
| 7      | c.444C > T | p.(=)           | No               | Absent           |
| 8      | c.614A > G | p.Tyr205Cys     | No               | Absent           |
| 9      | c.760G > A | p.Gly254Arg     | No               | Absent           |
| 10     | c.832G > T | p.Gly278Trp     | No               | Absent           |
| 11     | c.875-8376G > T | p.?              | No               | Absent           |
| 12     | c.921 T > C | p.(=)           | No               | Absent           |
| 13     | c.1082C > G | p.Ser361Cys     | No               | Absent           |
| 14     | c.1362G > C | p.Glu454Asp     | No               | Absent           |
| 15     | c.2026 + 650G > C | p.?              | No               | Absent           |
| 16     | c.2704G > T | p.Gly902Cys     | No               | Absent           |
| 17     | c.2771G > T | p.Gly924Val     | No               | Absent           |
| 18     | c.3184-1246A > T | p.?              | No               | Absent           |
| 19,20  | c.3376 T > C | p.Cys1126Arg    | No               | Absent           |
| 21     | c.3424A > G | p.Thr1142Ala    | No               | Absent           |
| 22     | c.3656-345G > T | p.?              | No               | Absent           |
| 23     | c.4391C > T | p.Ala1464Val    | No               | Absent           |
| 24     | c.5360G > A | p.Arg1787Gln    | No               | Absent           |

Note: http://cancer.sanger.ac.uk.

*ahttps://gnomad.broadinstitute.org.*
pathways (Che et al., 2016; Huang et al., 2019; Lu et al., 2016; Morell et al., 2013; Zhu et al., 2017). Some studies suggest that Notch acts as a tumor suppressor by inhibiting HCC proliferation (Huang et al., 2019; Lu et al., 2016; Qi et al., 2003; Viatour et al., 2011; Wang et al., 2009). On the other hand, in vitro studies using HCC cell lines suggest that Notch activation promotes HCC development based on the observation of an upregulation in progenitor cell differentiation, which has led some to propose Notch inhibition as a cancer therapy (Lu et al., 2016; Morell et al., 2013; Zhu et al., 2017). Conflicting findings warrant further examination of the downstream impacts of Notch dysfunction, as the mechanism of its involvement in HCC is likely complex.

Insertional mutagenesis studies support the specification of mouse analog Jag1 as a cancer-causing gene, however, Jag1 is not an established cancer gene in humans (Tate et al., 2019). Regardless, Jag1 variants have been described in human tumor tissue and its potential involvement in cancer cannot be ruled out. Our query of somatic Jag1 variants across cancer types indicates a low degree of tissue specificity; therefore, Jag1 could be implicated in the development of multiple cancer types, including HCC. Of note, the same Jag1 variant described in the proband (c.693_694del; p.Arg231Serfs*10) has been recorded in a sample of urinary tract carcinoma, supporting its possible role in cancerogenesis (Tate et al., 2019). Similarly, the missense Jag1 variant c.551G > A (p.Arg184His) reported in Patient 17 (Table 3) has also been recorded in adenocarcinoma of the large intestine (Tate et al., 2019). Of the Jag1 variants described in human cancer, missense variants occurred at the highest frequency (44.9%; Table S1). In contrast, the most common ALGS-causing pathogenic Jag1 variants are protein-truncating, including frameshift, nonsense, exon deletions, and splice site mutations. Only 15% of the Jag1 variants described in ALGS are missense variants (Gilbert et al., 2019). Interestingly, missense variants appear to be more common in cancer, but less common in ALGS. This observation could explain why HCC is a rare finding in ALGS.

No somatic Jag1 or Notch2 variants described in HCC have also been identified as germline in individuals with ALGS, and the majority of these Jag1 variants (4/6) result in deep intronic or synonymous protein changes that are unlikely to be pathogenic for ALGS, which is caused by gene haploinsufficiency (Table 5). Nine of the recorded somatic Notch2 variants in HCC were deep intronic, two resulted in synonymous protein changes, and 14 were rare missense variants (Table 6). Whether these somatic variants could be causative for ALGS is unclear, as only 19 Notch2 variants have been described in ALGS to date and the mechanism of pathogenesis is unclear. Moreover, germline pathogenic variants identified in ALGS cause abnormalities to arise during development in all tissues, in contrast to later-arising somatic variants that are confined to specific tissues. The Jag1 and Notch2 variants reported in the COSMIC database are somatic changes identified in tumor tissue, which could explain a lack of systemic ALGS as well as why the types of Jag1 and Notch2 variants observed in cancer differ significantly from those seen in ALGS.

Although in this report we are specifically focused on HCC, we also reviewed the frequency of Jag1 and Notch2 variants in hepatocellular adenoma and biliary tract carcinoma recorded in the COSMIC database. Pacheco et al. describe the first recorded Notch2 variant in an individual with ALGS and hepatocellular adenoma (p.C286S), however, this appears to be a variant of uncertain significance for ALGS, thus the disease association is weak (Pacheco, Monroe, & Horslen, 2018). There are 26 and 36 biliary tract carcinoma samples with Jag1 or Notch2 variants, respectively, reported in COSMIC (n = 3,069 human tumor samples). The majority of these variants occur at a high population frequency, are deep intronic variants, or else have not been described in ALGS. Thus, there is little evidence to suggest an association between Jag1 and Notch2 variants identified in hepatocellular adenoma or biliary tract carcinoma and ALGS, and the occurrence of liver cancer in individuals with ALGS appears to be confined to HCC, as we and others have now reported.

Among ALGS patients who present with severe liver manifestations, close monitoring of liver enzymes, tumor-marker α-fetoprotein, and periodic ultrasound and CT scan or MRI is standard protocol (Bhadri et al., 2005; Kaufman et al., 1987; Kim et al., 2005; Schwarzenberg et al., 1992; Tsai et al., 2010). However, no screening protocol currently exists for patients with attenuated ALGS phenotypes (Tsai et al., 2010). In the proband described, lack of severe hepatic symptoms prior to developing HCC precluded periodic cancer screening that could have enabled detection early enough for resection or transplantation to be viable. Keeffe et al. (1993) describe a similar case of a 46-year-old man with mild ALGS and no hepatic symptoms who developed HCC (Patient 28, Table 4). Like our proband, he was asymptomatic and was not diagnosed with ALGS until after the diagnosis of his more severely affected child. HCC was identified upon his initial physical examination, and although he underwent a successful LT, he died of recurrent HCC 14 months later. Given that AFP levels in our proband remained within a normal range during the course of her illness, its reliability as a tumor marker is questionable, and indeed we report eight cases of HCC in ALGS in which AFP levels stayed within a normal range (D’Souza et al., 2018; Galvez et al., 2020; Pham et al., 2015; Schwarzenberg et al., 1992; Valamparampil et al., 2020; Weiss et al., 2018). The development of HCC in two adults with mild ALGS and no liver involvement identifies an unmet need for increased surveillance of all individuals with an ALGS-associated Jag1 or Notch2 mutation, and also underscores the importance of genetic screening of family members to identify those who may be asymptomatic carriers of a pathogenic variant.

Conversely, it may also be prudent to consider the diagnosis of ALGS in patients presenting with HCC. The prevalence of HCC in the United States is roughly 7.7 cases per 100,000, with the prevalence of ALGS estimated at roughly one in 30,000 (Rich, Yopp, Singal, & Murphy, 2020; Saleh et al., 2016). It is well documented that ALGS is under-diagnosed in individuals with mild or sub-clinical symptoms and it is possible that some of these undiagnosed adults may be contributing to the prevalence of apparently isolated HCC, as we see in our proband and in Patient 28 (Saleh et al., 2016). Given the likelihood of other similar cases of HCC in adult asymptomatic ALGS that go unreported, it is possible that the incidence is significantly higher than has been predicted from case reports. Additionally, Wegmann et al. describe a patient with HCC who was not diagnosed with ALGS until
a postmortem evaluation of the medical history revealed characteristic clinical features (Wegmann et al., 1996). These cases suggest that ALGS should be considered in the differential diagnosis for individuals presenting with HCC, especially in those without a concomitant underlying risk factor or with a suspected or confirmed family history of ALGS, as this would change counseling.

To the best of our knowledge, only 35 cases of HCC in ALGS have been reported to date, including our proband. None of these cases address potential underlying risk factors for HCC, such as NAFLD, HBV or HCV infection, obesity, or alcoholism (Desai et al., 2019). Our research was limited by a lack of correlated genetic data, with 12 cases (36.3%) published prior to the discovery of JAG1 as the disease-causing gene in 1997 (Li et al., 1997). Even in reports published after 1997, the majority of diagnoses were based solely on clinical manifestations without molecular genetic confirmation. Consequently, there is no genotype–phenotype correlation between type of mutation and HCC risk. Finally, none of the cases presented address possible somatic mosaicism. Mosaicism of ALGS-causing variants could be associated with a milder phenotype and may explain the observation of HCC in adults with milder ALGS phenotypes (Giannakidis et al., 2001). Future sequencing studies should focus on uncovering possible molecular etiologies of HCC among individuals with ALGS in order to help affected individuals understand their risk and guide the clinical management of ALGS. Additionally, care should be taken when screening family members of those with ALGS for HCC, with an emphasis on imaging as a diagnostic tool over AFP, to ensure that this at-risk population is appropriately screened.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
David A. Piccoli, Ian D. Krantz, and Kathleen M. Loomes reviewed the patient’s medical records. Emma A. Schindler, Ian D. Krantz, and Melissa A. Gilbert conducted the literature review. Emma A. Schindler and Ian D. Krantz drafted the initial manuscript and Emma A. Schindler, Kathleen M. Loomes, and Melissa A. Gilbert edited the draft based on recommendations from co-authors and reviewers. Nancy B. Spinner performed genomic analysis for the proband and affected family members. All authors read and approved the final version of the manuscript.

DATA AVAILABILITY STATEMENT
Data sharing is not applicable to this article as no new data were created or analyzed in this study.

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