### ARTICLE





# The histologic presentation of hepatitis E reflects patients' immune status and pre-existing liver condition

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### Abstract

Infection with the hepatitis E virus (HEV) is one of the main causes of acute hepatitis worldwide. Given that, the histopathology of hepatitis E is relatively poorly characterized, and it is unclear what exactly determines its remarkable variability. The aim of our study was a systematic analysis of hepatitis E histology, especially with regard to the clinical setting, Fifty-two liver samples (48 biopsies, 1 liver explant, 3 autopsy livers) from 41 patients with molecularly proven hepatitis E (28 HEV genotype (gt) 3, three gt 1, one gt 4 and 9 undetermined gt) were systematically evaluated for 33 histopathologic features. Following one approach, the biopsies were assigned to one of five generic histologic patterns. In another approach, they were subjected to hierarchical clustering. We found that 23/41 (56%) patients were immunocompromised, whereas 18 (44%) had no known immunosuppression. Five patients (12%) had pre-existing liver disease (LD). The histopathologic spectrum ranged from almost normal to acute, chronic, and steato-hepatitis to subtotal necrosis, and was thus distributed across all five generic patterns. Hierarchical clustering, however, identified three histopathologic clusters (C1–C3), which segregated along the immune status and pre-existing LD: C1 comprised mostly patients with pre-existing LD; histology mainly reflected the respective LD without pointing to the additional hepatitis E. C2 comprised mostly immunocompetent patients; histology mainly displayed florid hepatitis. C3 comprised mostly immunocompromised patients; histology mainly displayed smoldering hepatitis. Accordingly, C1-C3 differed markedly with respect to their clinical and histopathologic differential diagnoses. Hierarchical clustering suggests three groups with distinct histopathologies, indicating biologically different manifestations of hepatitis E. The association of histopathologic changes with the patient's immune status and pre-existing LD plausibly explains the diversity of hepatitis E histopathology, and suggests that these factors are the crucial underlying determinants. We expect our results to improve patient management by guiding the clinico-pathologic diagnosis of hepatitis E.

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# Introduction

Hepatitis E is caused by a water- or food-borne, enterically transmitted infection with the hepatitis E virus (HEV), a non-enveloped, positive-strand RNA virus [1, 2]. HEV

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infection is one of the most common causes of acute hepatitis worldwide and thus represents a major global health problem [3, 4]. The disease shows clear differences between resource-rich and resource-limited countries in terms of viral genotypes (gt), mode of transmission and course of disease. Resource-limited countries are dominated by gt 1 and gt 2, which occur endemically as well as epidemically, are transmitted from humans to humans predominantly by contaminated drinking water and occasionally cause large epidemic outbreaks. By contrast, resource-rich countries are dominated by zoonotic transmission of HEV gt 3, which occurs worldwide, as well as gt 4, which is mainly found in China and Southeast Asia. Transmission of gt 3 and gt 4 often occurs through contaminated meat products [5].

The clinical course of hepatitis E is highly variable. HEV gt 3 infection is usually asymptomatic or subclinical but can also present as acute, self-limiting hepatitis, or even as acute-on-chronic liver failure in patients with pre-existing liver disease [3, 6]. Poor outcomes have also been described in pregnant women infected with HEV gt 1 [3, 6]. Originally regarded as an exclusively acute infection, the experience of the last decade has shown that hepatitis E of gt 3 can also develop a chronic course under immunosuppression [7].

Shortly after a peculiar form of epidemic hepatitis, later on designated as hepatitis E, had been identified as a distinct disease in India in the mid-fifties [8], the histologic changes observed in >70 cases of this outbreak were described [9]. For a long time, these reports, limited to the epidemic form of hepatitis E and enriched for severe courses, were the only ones available. Several decades later, the first histologic descriptions of autochthonous hepatitis E were published in smaller series [10-13], including HEV infection taking a chronic course [7, 14–16]. Although a comprehensive, comparative analysis of histology, especially with regard to characteristics of the host, is still lacking, from the body of literature on hepatitis E histology it is obvious that findings are highly variable and overlapping with other causes of hepatitis, thus, making a timely and accurate histopathologic diagnosis of HEV infection challenging [10, 14, 17–19]. Since the histologic changes reflect the interaction between the virus on the one hand and the patient on the other, we here have aimed at a systematic analysis of hepatitis E histology and its association with clinical parameters.

# Materials and methods

# Patients and liver tissue samples

Included into this study were 52 liver specimens (48 biopsies, 1 liver explant, and 3 autopsy livers) which have been

analyzed at the Department of Pathology and Molecular Pathology, University Hospital Zurich (USZ) between 2012 and 2018. The cohort comprised cases primarily submitted to the USZ and cases referred to us from other centers, including the Cantonal Hospitals St. Gallen and Lucerne, the University Hospitals Lausanne, Geneva, Basel and Bern, all Switzerland, the University Hospital Regensburg, Germany, and the Massachusetts General Hospital, Boston, MA, USA. For this study, all liver specimens submitted to our center were included from patients who had molecularly confirmed HEV infection (n = 41), i.e., having a positive HEV RNA PCR in blood and/or tissue. Two or more specimens taken at different time points were analyzed from eight of the 41 patients. Information on demographic, clinical, and virological parameters, including age, sex, immune status, pre-existing LD, and viral genotype, was retrieved retrospectively from the patient charts. This study was approved by the internal review board of the USZ and the Cantonal Ethics Committee of Zurich, Switzerland (KEK-ZH-Nr. 2013-0504).

#### Histopathologic evaluation of liver biopsies

All liver specimens were processed according to standard histologic methods. Stains included hematoxylin & eosin (H&E) and a connective tissue stain (such as Sirius red or Masson trichrome stain) for all cases, as well as an additional periodic acid-Schiff stain after diastase digestion (PAS-D). Histologic slides were re-evaluated independently by three experienced liver pathologists (DL, CS, AW). For all biopsies, the following 33 histopathologic features were systematically recorded and graded: portal inflammation (0-3), portal lymphocytes (0-2), portal plasma cells (0-2), portal histiocytes (0-2), portal ceroid-laden histiocytes (0-2), portal neutrophils (0-2), portal eosinophils (0-2), portal edema (no/yes), bile duct damage (0-2), ductular reaction (no/yes), interface activity (0-3), lobular inflammation (0-3), hepatocyte necrosis (0-3), lobular ceroidladen histiocytes (0-2), Kupffer cell activation (0-2), lobular disarray (0-2), rosettes (0-3), anisonucleosis (no/yes), hepatocyte damage (0-2), Mallory-Denk hyaline (no/yes), sinusoidal dilatation (no/yes), cholate stasis (no/yes), bilirubinostasis (0-2), steatosis microvesicular (0-2), steatosis macrovesicular (no/yes), Brunt score for steatosis (0-3) [20], fibrosis (0-3), sinusoidal fibrosis (no/yes), activity according to the classifications METAVIR (A0-A3) [21], and Batts-Ludwig (A0-A4) [22] as well as fibrosis according to the classifications METAVIR (F0-F4) [21], Batts-Ludwig (F0-F4) [22], and Ishak (0-6) [23] (Supplementary Table 1).

According to this evaluation, each specimen was assigned to one of the following five generic histologic patterns by all three pathologists: (1) Minimal active hepatitis, (2)

Table 1 Cohort characteristics: p	patients and li	ver specimens
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Patients, n	41
Median age at first biopsy (range)	56 (10–76)
Sex male/female, n (%)	27/14 (66%/34%)
HEV genotype (gt), $n$ (%)	- gt 1: 3 (7.3%)
	- gt 3: 28 (68.3%)
	- gt 4: 1 (2.4%)
	- NA: 9 (22.0%)
Immune status	
- Immunocompetent, n (%)	18/41 (44%)
$\bigcirc$ No known pre-existing LD, $n$ (%)	- 13/41 (32%)
$\bigcirc$ Pre-existing LD, <i>n</i> (%)	- 5/41 (12%)
- cirrhosis on ASH	O 3
- cirrhosis on NASH	O 1
- cirrhosis on ASH/NASH	O 1
- Immunocompromised, n (%)	23/41 (56%)
○ TPL (solid organs or HSC), n (%)	- 14/41 (34%)
- Liver, n	07
- Kidney, n	O 2
- Liver and kidney, n	O 1
- HSC, <i>n</i>	O 4
$\bigcirc$ Otherwise, <i>n</i> (%)	- 9/41 (22%)
- RA (methotrexate ± infliximab)	O 2
- MS (fingolimod)	O 2
- SLE and cryoglobulinemia (prednisolone and leflunomide)	O 1
- AIH (prednisone or prednisolone and tacrolimus)	O 2
- CLL (rituximab, leflunomide)	O 1
- Histiocytosis X (chemotherapy NOS)	O 1
Liver specimens, n	52
- Biopsy, n (%)	- 48/52 (92%)
- Explant liver, n (%)	- 1/52 (2%)
- Autopsy liver, n (%)	- 3/52 (6%)

*AIH* autoimmune hepatitis, *ASH* alcoholic steatohepatitis, *CLL* chronic lymphocytic leukemia, *gt* genotype, *HEV* hepatitis E virus, *HSC* hematopoietic stem cells, *LD* liver disease, *MS* multiple sclerosis, *NA* not available, *NASH* nonalcoholic steatohepatitis, *NOS* not otherwise specified, *RA* rheumatoid arthritis, *SLE* systemic lupus erythematosus, *TPL* transplantation.

Predominantly lobular hepatitis, (3) Predominantly portal hepatitis, (4) Steatohepatitis, and (5) Extensive necrosis (pattern definition see Supplementary Table 2). In case of discrepancies, consensus was found in a panel discussion.

# Statistics

To quantify the inter-rater agreement of the assessments of the histopathologic features, the weighted kappa statistic was computed using quadratic weights. For the histologic pattern variables, the usual kappa statistic was used [24]. Divisive hierarchical clustering of the 33 histopathologic features was conducted according to the algorithm by Kaufman and Rousseeuw [25]. Since the histopathologic features were on ordinal scales with different ranges and a few values were missing, an extension of Gower's generalized dissimilarity coefficient to compute dissimilarities for the clustering procedures was used (with the algorithm "daisy" available in the R-package "cluster") [25]. Gower's method also allows for different weighting of features. We assigned a weight of 1 to all features, except the following: Steatosis macrovesicular and Brunt-Score for steatosis received a weight of 1/2 each, activity METAVIR and activity Batts-Ludwig received a weight of 1/2 each, and fibrosis METAVIR, fibrosis Batts-Ludwig, and fibrosis Ishak received a weight of 1/3 each. This was done to avoid overweighting these features in the dissimilarity calculation as they are measures of the same histologic parameters [25]. Finally, the results of the cluster analysis were then displayed in a heatmap, in which the histopathologic features have been properly scaled such that criteria with different number of categories can be compared. All statistical analyses were performed in the R programming language [26] using base packages, as well as the psy [27], cluster [28], gplots [29], and vcd [30] packages.

### Results

# The cohort comprises mostly HEV genotype 3 infections in immunocompetent as well as immunocompromised individuals

Basic demographic and clinical characteristics as well as the results of genotyping were retrospectively retrieved from the charts of all 41 patients. These are summarized in Table 1 and listed in more detail in Table 2. From eight patients, there were two or more liver samples available over time, yielding a total of 52 liver specimens for analysis. The age at the first liver biopsy ranged from 10 to 76 years, with a median age of 56 years. Twenty-seven patients (66%) were male and 14 (34%) were female. HEV infection was proven molecularly by positive HEV RNA PCR in blood and/or tissue in all 41 patients. In 32 of these 41 patients, the HEV gt was determined, with the following distribution: gt 3 in 28 patients (68.3%), gt 1 in three patients (7.3%), and gt 4 in one patient (2.4%). For nine patients (22.0%), no information on the HEV genotype was available.

Four different patient groups were represented based on clinical considerations [3, 4]. These comprised immunocompetent individuals without or with known pre-existing

Table 2 Fifty-two liver specimens from 41 patients.	2 Fitt		T	-										
Pat. No.	Sex	Age (y)	) Sample No.	Liver specimen	Histologic pattern	Histo- pathologic cluster	Immune status				HEV geno- type	Clinical presentation	Outcome with respect to HEV infection	Previously reported
							Immunocompetent	ant	Immunocompromised	mpromised	;			
							No known F pre- existing LD	Pre-existing LD	TPL	other	I			
Pat.01	в	35	la	Biopsy 2 months post TPL	Predominantly portal hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 3 months	Spontaneous resolution	e
		35	lb	hs	Predominantly portal hepatitis	Cluster 3								B
Pat.02	Е	56	2a	Biopsy 7.5 months post TPL	Minimal active hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 3.5 years	Viral clearance after reduction of immunosuppressive drug	a, f
		57	2b	Biopsy 1 year 5.5 months post TPL	Minimal active hepatitis	Cluster 3							treatment and ribavirin treatment	e
		57	2c	Biopsy 1 year 10 months post TPL	Predominantly portal hepatitis	Cluster 3								a
		58	2d	Biopsy 2 years 2 months post TPL	Predominantly portal hepatitis	Cluster 2								a, f
		58	2e	SL	Predominantly portal hepatitis	Cluster 2								e
Pat.03	f	29	3a	Biopsy 4 months post TPL	Predominantly lobular hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 5 months	Spontaneous resolution	ę
		30	3b	Biopsy 7 months post TPL	Predominantly portal hepatitis	Cluster 3								5
Pat.04	f	33	4	Biopsy	Predominantly portal hepatitis	Cluster 3			Liver & kidney		gt 3	Prolonged elevation of transaminases for 1 year 10 months	Viral clearance after reduction of immunosuppressive drug treatment	p
Pat.05	f	22	S	Biopsy	Predominantly lobular hepatitis	Cluster 2	X				gt 1	Acute hepatitis after travel to India	Spontaneous resolution	q
Pat.06	Ξ	67	9	Biopsy	Predominantly portal hepatitis	Cluster 1	X				gt 3	Acute hepatitis	Spontaneous resolution	
Pat.07	E	48	7	Biopsy	Predominantly portal hepatitis	Cluster 3			Kidney		gt 3	Prolonged elevation of transaminases for >6 months	Viral clearance after reduction of immunosuppressive drug treatment	q
Pat.08	Е	59	×	Biopsy	Predominantly lobular hepatitis	Cluster 2	×				gt 4	Acute hepatitis after travel to Corsica and meal in an Asia restaurant	Spontaneous resolution	Ą
Pat.09	f	26	6	Liver explant	Extensive necrosis	Cluster 2	Х				gt 1	Acute hepatitis after Liver failure travel to India requiring TP	Liver failure requiring TPL	b, d, e, f

Pat.10 m 36 10   Pat.11 m 41 11   Pat.12 m 45 12   Pat.13 m 52 12   Pat.14 m 31 14   Pat.15 m 66 12   Pat.16 m 50 16   Pat.16 m 50 16   Pat.13 f 66 12   Pat.16 m 50 16   Pat.17 f 67 17   Pat.18 f 50 19	6 5 7 3 2 1 0		Predominantly lobular hepatitis Predominantly portal hepatitis Predominantly portal hepatitis Predominantly	Cluster 3	Immunocompetent No known Pre pre-	stent Pre-existing LD	Immunoco	Immunocompromised				
m 36 m 41 m 45 m 32 m 31 f 67 f 50 f 50	6 IS 4 3 12 II 0		<u>s</u>	Cluster 3	No known pre-	Pre-existing LD						
m 36 m 41 m 45 m 52 m 31 m 66 m 50 f 67 f 50	10 0 13 13 13 10 16		S	Cluster 3	existing LD		TPL	other				
m 41 m 45 m 52 m 31 m 66 m 50 f 67 f 50	1				х				gt 3	Acute hepatitis	Spontaneous resolution	Ą
m 45 m 52 m 31 m 66 m 66 f 67 f 50	12 IS 14 IS 10			Cluster 2	x				gt 3	Acute hepatitis	Spontaneous resolution	q
m 52 m 31 m 66 m 50 f 67 f 50	16 IS [4 3			Cluster 2	x				gt 3	Acute hepatitis	Spontaneous resolution	p
m 31 m 66 m 50 f 67 f 50	6 IS 4		hortan mepanan	Cluster 2	x				gt 3	Acute hepatitis	Spontaneous resolution	d, e
m 66 m 50 f 67 f 50	رہ ب <del>ک</del>		Predominantly portal hepatitis	Cluster 3			Kidney		gt 3	Prolonged elevation of transaminases for 1 year 10 months	Viral clearance after reduction of immunosuppressive drug treatment and ribavirin treatment	q
m 50 f 67 f 50	16	Biopsy 3 J years post TPL	Predominantly portal hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 3 months	Viral clearance after reduction of immunosuppressive drug treatment - ribavirin treatment	d, e
f 67 f 50		Biopsy	Predominantly lobular hepatitis	Cluster 3	×				gt 1	Acute hepatitis, Indian citizen traveling to USA	Spontaneous resolution	c, d, f
f 50	17	Biopsy	Predominantly portal hepatitis	Cluster 3	X				gt 3	Acute hepatitis	Spontaneous resolution	þ
	18	Biopsy	Predominantly lobular hepatitis	Cluster 3				SLE (prednisolone, leftunomide)	gt 3	Acute hepatitis	Spontaneous resolution	q
Pat.19 m 74 19	19	Biopsy 21 days prior to death	Steatohepatitis	Cluster 1		Cirrhosis on ASH			NA	Acute-on- chronic LD	Liver failure, death	q
Pat.20 m 52 20	20	Biopsy	Predominantly lobular hepatitis	Cluster 3				MS (fingolimod)	NA	Acute hepatitis	Resolution after stop fingolimod	p
1 73	21	Biopsy	Steatohepatitis	Cluster 2	x				gt 3	Acute hepatitis	Spontaneous resolution	q
f 65	22	Biopsy	Predominantly portal hepatitis	Cluster 3				CLL (rituximab, lenalidomide)		Prolonged elevation of transaminases for 4 years	No viral clearance, even infer reduction of immunosuppressive drug treatment and ribsvirin treatment - death (not related to HEV infection)	p
f 62	23			Cluster 3				RA (methotrexate)	gt 3	Acute hepatitis	Spontaneous resolution	p
Pat.24 m 70 24	24	Biopsy	Predominantly portal hepatitis	Cluster 2	X				gt 3	Acute hepatitis	Spontaneous resolution	þ
Pat.25 m 20 24	25	Biopsy	Predominantly lobular hepatitis	Cluster 3				Histiocytosis X (CHTx)	gt 3	Acute hepatitis	Spontaneous resolution	p

Table 2 (continued)

**SPRINGER NATURE** 

Pat. No.														
		Sex Age (y)	) Sample No.	Liver specimen	Histologic pattern	Histo- pathologic cluster	Immune status	SI			HEV geno- type	Clinical presentation	Outcome with respect to HEV infection	Previously reported
							Immunocompetent	petent	Immunoc	Immunocompromised				
							No known pre- existing LD	Pre-existing LD	TPL	other				
Pat.26	в	60	26	Biopsy	Minimal active hepatitis	Cluster 3			HSC		gt 3	Prolonged elevation of transaminases for 6 months	No viral clearance, death (not related to HEV infection)	p
Pat.27	f	66	27a	Biopsy 10 days prior to death	Steatohepatitis	Cluster 1		Cirrhosis on ASH/ NASH			gt 3	Acute-on- chronic LD	Liver failure, death	f
		99	27b	Autopsy	Extensive necrosis	Cluster 1								
Pat.28	в	71	28	Biopsy	Predominantly lobular hepatitis	Cluster 2	×				NA	Acute hepatitis	Spontaneous resolution	
Pat.29	f	67	29	Biopsy	Predominantly lobular hepatitis	Cluster 2				RA (methotrexate, infliximab)	NA	Acute hepatitis	Spontaneous resolution	f
Pat.30	В	64	30	Biopsy	Predominantly portal hepatitis	Cluster 2				AIH (prednisone)	gt 3	Acute hepatitis	Spontaneous resolution	
Pat.31	В	76	31a	Biopsy 8 days prior to death	Steatohepatitis	Cluster 1		Cirrhosis on ASH			gt 3	Acute-on- chronic LD	Liver failure, death	
		76	31b	Autopsy	Extensive necrosis	Cluster 1								
Pat.32	н	65	32a	Biopsy 3 years post TPL	Predominantly portal hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 3 years		
		67	32b	Biopsy 6 years post TPL	Predominantly portal hepatitis	Cluster 1							treatment and ribavirin treatment	
Pat.33	f	61	33	Biopsy	Predominantly portal hepatitis	Cluster 3			HSC		gt 3	Acute hepatitis	Spontaneous resolution	
Pat.34	В	61	34	Biopsy	Predominantly portal hepatitis	Cluster 2			HSC		NA	Prolonged elevation of transaminases for 9 months	No viral clearance, death (not related to HEV infection)	
Pat.35	Ш	59	35a	Biopsy 4 days prior to death	Steatohepatitis	Cluster 1		Cirrhosis on NASH			NA	Acute-on- chronic LD	Liver failure, death	
		59	35b	Autopsy	Extensive necrosis	Cluster 1								
Pat.36	f	45	36	Biopsy	Predominantly portal hepatitis	Cluster 2				MS (fingolimod)	NA	Acute hepatitis	Spontaneous resolution	
Pat.37	н	50	37	Biopsy	Predominantly lobular hepatitis	Cluster 3			HSC		NA	Acute hepatitis	Spontaneous resolution	
Pat.38	Ъ.	13	38	Biopsy	Minimal active hepatitis	Cluster 3				AIH (tacrolimus, prednisolone)	gt 3	Prolonged elevation of transaminases for 7.5 months	Viral clearance after reduction of immunosuppressive drug treatment and ribavirin treatment	

	1000) 4													
Pat. No	. Sex	Age (y)	Pat. No. Sex Age (y) Sample No. Liver specie	Liver specimen	Histologic pattern	Histo- pathologic cluster	Immune status	s			HEV geno- type	Clinical presentation	Outcome with respect to Previously HEV infection reported	Previously reported
							Immunocompetent	etent	Immunocc	Immunocompromised				
							No known pre- existing LD	No known Pre-existing LD pre- existing LD	TPL	other				
Pat.39	в	58	39	Biopsy	Predominantly portal hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation Viral clearance after of transaminases for ribavirin treatment 7 months	Viral clearance after ribavirin treatment	
Pat.40	Ξ	10	40a	Biopsy	Minimal active hepatitis	Cluster 3			Liver		gt 3	Prolonged elevation of transaminases for 2.5 years	Prolonged elevation Viral clearance of transaminases for after reduction of 2.5 years immunosuppressive drug treatment	
		12	40b	Biopsy	Predominantly portal hepatitis	Cluster 3								
Pat.41	f	51	41	Biopsy 17 days prior to death	Steatohepatitis	Cluster 1		Cirrhosis on ASH			NA	Acute-on- chronic LD	Liver failure, death	
1114		-	110 4	1-1-1-		1 1 1								

AIH autoimmune hepatitis, ASH alcoholic steatohepatitis, CHTx chemotherapy, CLL chronic lymphocytic leukemia, d days, gt genotype, HSC hematopoietic stem cells, LD liver disease, mo months, MS multiple sclerosis, NA not available, NASH nonalcoholic steatohepatitis, RA rheumatoid arthritis, SLE systemic lupus erythematosus, TPL transplantation, y years.

<sup>a</sup>Protzer U et al. [14] Mod Pathol 2015;28(4):523–32. <sup>b</sup>Chijioke O et al. [18] Front Med 2015;30;2:20.

<sup>c</sup>Friedman LS et al. [42] N Engl J Med 2016;375:2082–92.

<sup>d</sup>Lenggenhager D, Gouttenoire J et al. [31] J Hepatol 2017;67:471–479.

<sup>e</sup>Lenggenhager D and Weber A [43] Gastroenterol Clin North Am 2017; 46:393-407.

<sup>f</sup>Lenggenhager D and Weber A [35] Adv Anat Pathol 2018;25:273-81.

LD and immunocompromised patients after transplantation (solid organ or hematopoietic stem cell (HSC)), or otherwise immunocompromised. With respect to these categories, 18 of the 41 patients were immunocompetent (43.9%), with a total of 21 liver specimens (21/52; 40.4%). Of these 18 patients, 13 did not have any known preexisting LD (13/41; 31.7%), with a total of 13 liver specimens (13/52; 25.0%), and five patients had a known pre-existing LD (5/41; 12.2%), with a total of eight liver specimens (8/52; 15.4%). All five patients with known preexisting LD had cirrhotic livers. Etiologies of the cirrhosis included ASH (3/5), NASH (1/5), or ASH/NASH (1/5). Twenty-tree patients were immunocompromised (23/41; 56.1%), with a total of 31 liver specimens (31/52; 59.6%). The group of immunocompromised patients comprised 14 patients after solid organ or HSC transplantation (14/41; 34.1%), with totally 22 liver specimens (22/52; 42.3%), and nine otherwise immunocompromised patients (9/41; 22.0%), with totally nine liver specimens (9/52; 17.3%). Of the 14 transplanted patients, seven received a liver (7/14), two a kidney (2/14), one a liver and a kidney (1/14), and four HSC (4/14). Patients immunocompromised for other reasons included treatment for rheumatologic/autoimmune diseases (i.e., rheumatoid arthritis, multiple sclerosis, systemic lupus erythematosus, autoimmune hepatitis) or neoplasia (i.e., chronic lymphocytic leukemia, histiocytosis X).

# A broad and highly variable spectrum of histopathologic findings can be observed in livers of patients infected with HEV

First, all 52 liver specimens were re-evaluated independently by three expert liver pathologists, each recording systematically 33 histopathologic features (Supplementary Table 1). Inter-rater agreement in this independent evaluation was between good to excellent, with kappa values between 0.4 and 0.9 (two-thirds ≥0.75) for most histopathologic features (Supplementary Fig. 1). The microscopic review revealed a wide range of histopathologic findings, which varied significantly from case to case. This spectrum of hepatitis E histology is illustrated in Supplementary Fig. 2a-e. To corroborate the HEV infection on a tissue level, immunohistochemistry was performed in cases with sufficient block material, which was possible in a total of 40 tissue samples. Various staining patterns were observed as previously described [31]. Remarkably, cases with only minimal lobular inflammatory activity usually showed the strongest and most widespread positivity for HEV ORF2 protein (Supplementary Fig. 2f).

Collectively, these findings demonstrate that on the level of their histopathologic manifestation, the hepatitis E cases of the cohort were highly variable.

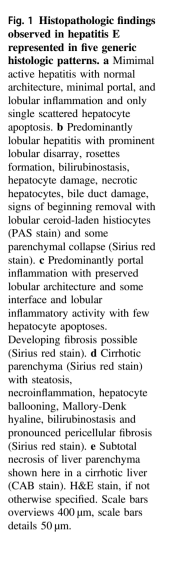
# Hepatitis E presents in five different generic histologic patterns

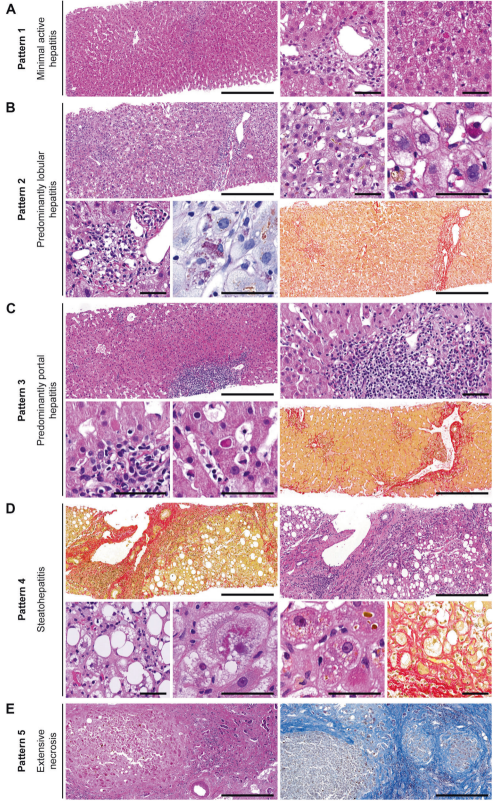
Next, we aimed to assign a generic histologic pattern to each liver specimen. Such patterns are considered to reflect the limited repertoire of the liver to respond to injuries and are widely used in routine histopathologic practice to get direction for the underlying etiology and correct diagnosis [32]. The cases of our cohort revealed the following distribution among the five histologic patterns (Fig. 1): (1) Minimal active hepatitis: 5/52 (9.6%; Fig. 1a), (2) Predominantly lobular hepatitis: 11/52 (21.2%; Figs. 1b), (3) Predominantly portal hepatitis: 26/52 (50.0%; Figs. 1c), (4) Steatohepatitis: 6/52 (11.5%; Fig. 1d) and (5) Extensive liver necrosis: 4/52 (7.7%; Fig. 1e). Thus, similar as observed for the individual histopathologic features, also the histologic patterns displayed a broad distribution among the cases of our cohort (Table 2 and Supplementary Fig. 3a). Again, the inter-rater agreement for those patterns was excellent (kappa value ≥0.75) (Supplementary Fig. 1). Of note, several of the histologic patterns were unequally associated with specific patient characteristics, some of them with statistical significance. For example, in the group of patients with pre-existing LD, the histologic patterns "steatohepatitis" and "extensive necrosis" were statistically significantly over-represented, whereas the pattern of "predominantly portal hepatitis" was statistically significantly under-represented (Supplementary Fig. 3b).

Collectively, these findings demonstrate that the histopathologic manifestation of hepatitis E covered a broad spectrum of well-established generic histologic patterns, some of which were preferentially associated with specific patient characteristics.

# Hierarchical clustering of 33 histopathologic features identifies three histopathologic clusters of hepatitis E

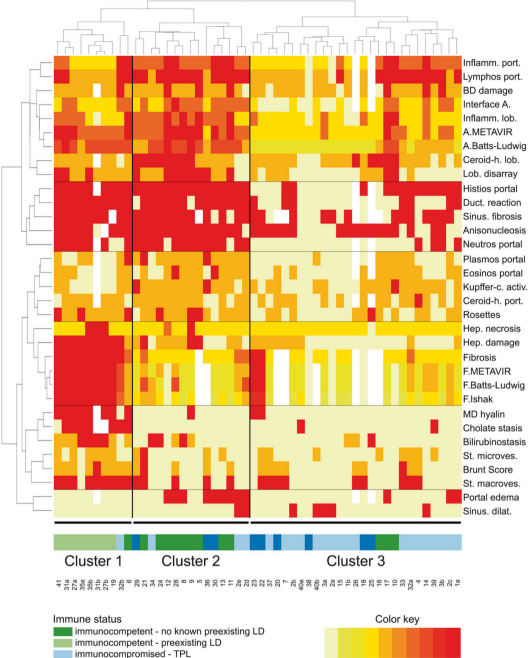
With the aim of uncovering any structure that may exist in this rich data set, regardless of any predetermined categories such as the patterns previously studied, we then decided to take a systematic approach by performing a cluster analysis. The principle of this analysis was to identify clusters of samples in which the samples within each cluster were histologically similar, but different from those in the other clusters. Specifically, we conducted hierarchical clustering of the samples with respect to the 33 histopathologic features recorded, as well as the other way around. Remarkably, this approach displayed an easy to identify structure in the data set, revealing three clusters of samples (C1–C3) with distinct histopathology. The liver samples were distributed as follows: Ten of the





52 samples (19.2%) in C1, 15 samples (28.8%) in C2 and 27 samples (52%) in C3 (Fig. 2). The 33 histopathologic features were clearly represented differently in the three

clusters. Of note, they were not randomly distributed, but clustered mainly according to the different pathologies they reflect (i.e., from top to bottom: features of acute



immunocompromised - otherwise

0 0.2 0.4 0.6 0.8 1

**Fig. 2 Hierarchical clustering of histopathologic findings.** Hierarchical clustering of the 52 liver specimens (x axis) with respect to the 33 histopathologic features (y axis) represented in a heatmap: Identification of three clusters (C1–C3) that segregate along the patient's immune status and pre-existing LD. The 33 histopathologic features clustered mainly according to the different pathologies they reflect (i.e., from top to bottom: features of acute inflammatory activity, features of late

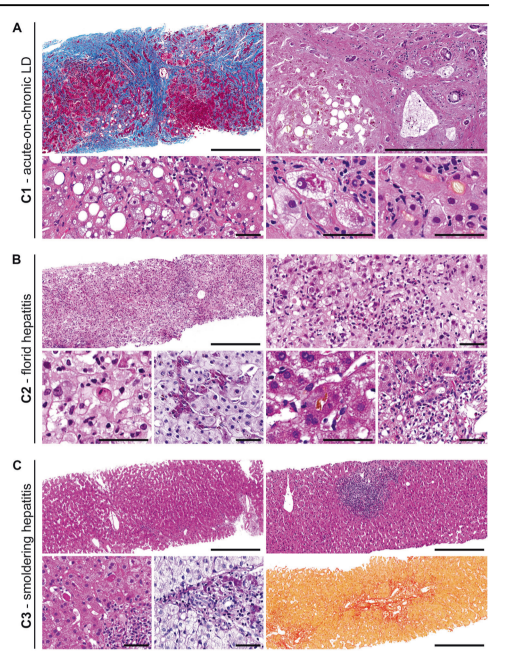
inflammatory activity, features of early subacute inflammatory activity, features of late subacute inflammatory activity, features of chronicity and features of cellular damage). Only two features (portal edema and sinusoidal cellular damage). Only two features (portal edema and sinusoidal dilatation) did not cluster in any of the above mentioned groups. A activity, BD bile duct, Ceroid-h. Ceroid-histiocytes, Duct. ductular, F fibrosis, Hep. hepatocyte, Inflamm. inflammation, Kupffer-c. activ. Kupffer cell activation, LD liver disease, lob. lobular, MD Mallory-Denk, port. portal, Sinus. sinusoidal, St. steatosis, TPL transplantation.

subacute inflammatory activity, features of chronicity and features of

dilatation) did not cluster in any of the above mentioned groups (Fig. 2).

Each of the three clusters was characterized by a predominance of different histopathologic features (Fig. 3):

### Fig. 3 Histopathologic findings observed in hepatitis E represented in three histopathologic clusters. a Samples in C1 showed mainly features of acute-on-chronic LD, such as cirrhosis (Masson trichrome stain), necrosis, steatosis, steatohepatitis, hepatocyte damage, Mallory-Denk hyaline and bilirubinostasis (from upper left to lower right). b Samples in C2 displayed mainly features of florid hepatitis, such as lobular inflammation and parenchymal disarray, foci of hepatocyte necrosis and many scattered apoptotic hepatocytes as well as signs of beginning removal with lobular ceroid-laden histiocytes (PAS-D stain), cholestasis and bile duct damage (from upper left to lower right). c Samples in C3 were mainly characterized by features of smoldering hepatitis. ranging from minimal histologic changes over predominantly portal based inflammation with mild interface and lobular activity with only rare apoptotic hepatocytes as well as signs of previous damage such as ceroidladen histiocytes (PAS-D stain) in portal tracts and in some cases with fibrosis (Sirius red stain) (from upper left to lower right). H&E stain, if not otherwise specified. Scale bars overviews 400 µm, scale bars details 50 µm. C cluster, LD liver disease.



Samples in C1 showed mainly features of acute-on-chronic LD, such as cirrhosis, necrosis, steatosis, steatohepatitis, hepatocyte damage, Mallory–Denk hyaline and cholestasis (Fig. 3a). Samples in C2 displayed mainly features of florid hepatitis, such as lobular inflammation, parenchymal disarray, foci of hepatocyte necrosis and many scattered apoptotic hepatocytes as well as signs of beginning removal with lobular ceroid-laden histiocytes, cholestasis and bile duct damage (Fig. 3b). Samples in C3 were mainly characterized by features of smoldering hepatitis, ranging from minimal histologic changes over mild to more pronounced portal based inflammation, mostly with mild interface and lobular activity with only rare apoptotic hepatocytes, as well

as signs of previous damage such as ceroid-laden histiocytes in portal tracts and in some cases with fibrosis (Fig. 3c).

Thus, similar as observed for the five histologic patterns, also the three clusters displayed a broad distribution among the cases of our cohort (Table 2 and Supplementary Fig. 4a), and were unequally associated with specific patient characteristics, some of them with statistical significance. For example, in C1 (acute-on-chronic LD) the group of patients with pre-existing LD was statistically significantly over-represented, whereas in C2 (florid hepatitis) the group of immunocompetent patients without pre-existing LD was statistically significantly over-represented. In contrast, in C3

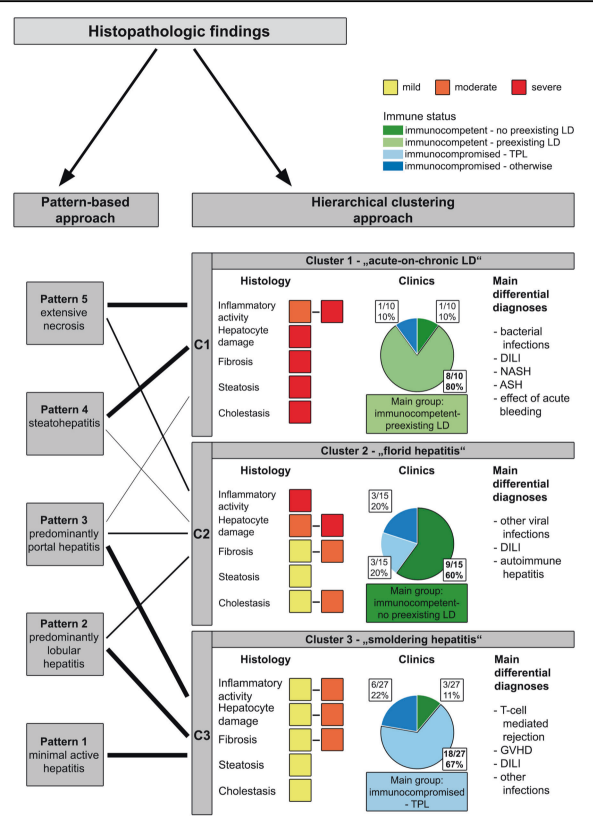


Fig. 4 Association of the three clusters with the clinical setting and the five histologic patterns. Clinico-pathologic settings of the three clusters including their most relevant differential diagnoses. The thickness of the lines between the patterns and the clusters represent

the relative association of both. ASH alcoholic steatohepatitis, C cluster, DILI drug-induced liver injury, GVHD graft-versus-host disease, LD liver disease, NASH nonalcoholic steatohepatitis, TPL transplantation.

(smoldering hepatitis), the group of immunocompetent patients without pre-existing LD was statistically significantly under-represented (Supplementary Fig. 4b).

Collectively, hierarchical clustering with respect to the 33 histopathologic features identified three histopathologic clusters, which showed distinct histopathologies, some of which were preferentially associated with specific patient characteristics.

# Three histopathologic clusters reflect different clinical scenarios

Next, we sought to investigate the association between the three clusters C1-C3 identified by statistical cluster analysis methods, and the patients' immune status and preexisting liver condition, considered to be relevant for the management of patients with hepatitis E [4]. Overall, it turned out that each of the three clusters C1-C3 was dominated by one main patient group (Fig. 4): C1 representing acute-on-chronic liver disease was dominated by patients with pre-existing LD (8/10; 80%); C2 representing florid hepatitis was dominated by immunocompetent patients without known pre-existing LD (9/15; 60%); C3 representing smoldering hepatitis was dominated by immunocompromised transplanted patients (18/27; 67%). Samples from patients immunocompromised for other reasons were found in all three clusters without a clear predominance. Of note, the five histologic patterns were represented differently in the three clusters (Fig. 4 and Supplementary Fig. 5a), and the three clusters were unequally associated with the five patterns, some of them with statistical significance. For example, the patterns "extensive necrosis" and "steatohepatitis" were statistically significantly over-represented in C1 (Supplementary Fig. 5b).

Moreover, we sought to investigate the association between the three clusters C1–C3 and the outcome with respect to the HEV infection. After 3 months as well as after 6 months, a strong positive association was found for C1 and "liver failure, death/liver failure, transplantation", and a strong negative association was found for C3 "liver failure, death/liver failure, transplantation." Furthermore, a strong positive association was found for C2 and "resolved infection", but only after 3 months (Table 2 and Supplementary Fig. 6).

Collectively, these findings not only show that the three histopathologic clusters are differentially associated with the five histologic patterns, but also suggest that they reflect different clinical scenarios, which seem to be defined by the patients' immune status and pre-existing liver condition, and also to affect the outcome of the HEV infection.

# Discussion

The findings of our study provide a rational explanation for the well-known broad and at first sight puzzling spectrum of histopathologic changes observed in hepatitis E. These results were made possible by (1) the availability of a significant number of hepatitis E tissue samples from patients with very different clinical backgrounds and (2) by pursuing a novel data-driven statistical approach. This allowed us to also conduct comparative studies on the different clinical settings of hepatitis E, which goes beyond the previous reports on narrowly defined cohorts such as hepatitis E in epidemic outbreaks or in patients with immunosuppression.

With both methods, the traditional pattern-based approach as well as the cluster analysis, we were able to structure the broad histopathologic spectrum of hepatitis E represented in this cohort. Of note, we ultimately found overlapping results between the two different approaches. Nevertheless, a strength of the data-driven approach is its independence of any predetermined category. We therefore consider the resulting clustering to be a solid basis for the classification of hepatitis E histopathology.

Of particular interest is the first cluster (C1, acute-onchronic liver disease), which includes mainly patients with pre-existing LD. From a histopathologic perspective, it is noteworthy that the observed changes were determined primarily by the respective pre-existing liver disease, i.e., cirrhosis on the background of ASH or NASH. Of note, beside cholestatic changes, considered to reflect hepatic decompensation, HEV infection in these cases showed no additional characteristic or even specific histologic features. As the samples in C1 mostly showed advanced fibrosis or cirrhosis as well as signs of steatohepatitis, the most important histopathologic differential diagnosis in this group is acute (non)alcoholic hepatitis. However, from a clinical perspective the differential diagnosis is much broader, including other reasons for acute decompensation such as bacterial infections, drug-induced liver injury (DILI), complications of acute bleeding, etc. Five of seven patients in C1 had cirrhosis and eventually died as a result of hepatitis E, in line with the previously reported poor prognosis in this patient group [6, 33]. A detailed clinical description of the patients with pre-existing LD who died in the context of HEV infection shall be reported elsewhere. Interestingly, the two patients in C1 who did not die (pat. 19 and 32), presented with only incomplete cirrhosis, presumably reflecting a higher reserve in coping with HEV infection. In conclusion for C1, knowledge and awareness of the specifics of (acute) hepatitis E in pre-existing liver disease is essential for both pathologists and clinicians. In daily practice, pronounced cholestatic changes and/or significantly elevated ALT values should prompt them to systematically look for reasons for liver decompensation, including hepatitis E [34].

The second cluster (C2, florid hepatitis) revealed an overrepresentation of immunocompetent patients without any known pre-existing LD, but included also five patients under immunosuppression. The fact that those five patients presented with high inflammatory activity may point to a relatively mild immunosuppression at the time of HEV infection. Histology was dominated by highly active hepatitis. The three top differential diagnoses from a pathological as well as from a clinical point of view are other viral infections, DILI, and autoimmune hepatitis [17, 35].

The third cluster (C3, smoldering hepatitis) included mainly patients under (probably more intense) immunosuppression. In line with an immune-mediated pathogenesis of hepatitis E, only mild to moderate inflammatory activity was observed in these patients. However, 13/22 patients in this group experienced a prolonged infection (3 months or longer, i.e., chronic hepatitis E), some of them also with developing fibrosis, which is in line with the current literature [7, 16]. In two of the immunocompromised patients in C3, histology revealed also cirrhotic changes. Patient 22 with chronic lymphocytic leukemia under rituximab/lenalidomid treatment had chronic hepatitis E persisting for 4 years. The etiology of liver cirrhosis in this patient was possibly a combination of hepatitis E and NASH, for which hints were found in histology. Patient 23 with a history of rheumatoid arthritis may have developed cirrhosis due to methotrexate treatment. It is possible that the three patients classified as immunocompetent in C3 (pat. 10, 16, and 17) had a hitherto unrecognized immunodeficiency, and as a consequence developed less active inflammation. Alternatively, since the biopsies showed clear signs of subacute inflammation, they may also have been taken at a later stage of HEV infection. The most important differential diagnoses from a pathologic as well as from a clinical point of view are other (viral) infections and DILI, as well as (acute) T-cell mediated rejection in patients with solid organ TPL and graft-versus-host disease in patients with HSC TPL [17, 35, 36].

Interestingly, in one and the same patient, the histologic changes caused by hepatitis E can vary considerably depending on the time the biopsy was taken, which most likely reflects the variable intensity of the patients' immunosuppression. For example, three biopsies from a liver transplant patient (pat. 2) clustered in C3 early during infection, while two biopsies taken at a later point in time were found in C2.

In routine diagnostics, both the pathologist and the clinician are not confronted with particular entities but with their manifestations, from which they must deduce the distinct etiologies behind them. The findings of our study suggest that for histopathologic—as well as clinical diagnosis of hepatitis E, it is advisable to consider the patients' immune status and pre-existing liver condition. This approach allows to sharpen and narrow down the differential diagnostic spectrum. For example, when being confronted with a little active pattern, hepatitis E should be among the top differential diagnoses in an immunocompromised patient, but it may be considered less likely in an immunocompetent individual.

Considering that the histopathologic changes in a tissue are a direct reflection of an organism's exposure to a harmful agent, such as an infectious particle, it is obvious that the histopathologic changes observed in relation to a particular disease are not only of interest to pathology but also provide insight into the biology of the disease, in particular with respect to immunologic aspects [37, 38]. We postulate that the associations of histologic patterns with the immune status and pre-existing liver condition of patients shown here reflect not only a random statistical association but have a causal link.

The cohort underlying this study is, to the best of our knowledge, the world's largest hepatitis E biopsy cohort comprising >50 specimens taken from patients with different immune and liver status, enriched with gt 3 infections [19]. With a predominance of HEV gt 3 and some travel associated gt 1 infections, it can be considered as representative not only for Europe, but also for North America where awareness of hepatitis E as a relevant health problem has increased recently [4, 39, 40]. Nevertheless, there are some potential limitations. First, despite the above mentioned high number of hepatitis E specimens, the sample size is still relatively low with respect to the applicability of confirmatory statistical analysis methods. Moreover, our results may not be generalizable to countries in which other gt predominate, such as Asia with gt 1 and gt 4 infections. However, it is conceivable that the histopathologic findings of cases of acuteon-chronic liver failure, as characterized here in C1, could be caused by gt 1, gt 2 or gt 4 infection as well [41]. Other potential limitations are the disproportionately high number of hepatitis E cases with a prolonged/chronic course in immunocompromised patients, since these are biopsied in a higher proportion, and the retrospective setting without standardized conditions, especially for the time point of the biopsy procurement during the infection.

In summary, the results of our study provide a plausible explanation for the broad, only at first glance unstructured histopathologic spectrum observed in hepatitis E, and provide insight into the underlying determinants. From these findings we expect not only a better understanding of the biology of HEV infection, but also guidance to the pathologic and clinical diagnosis, which should improve the care of patients with hepatitis E. Acknowledgements We would like to thank Christine Mittmann, André Fitsche as well as Martina Storz and her team for their excellent technical assistance.

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# **Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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