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販売名(企業名)	別紙のとおり								
研究報告の概要	<p>問題点：齧歯類の重症疾患の原因ウイルスとされていた cardiovirus がヒトにおいても存在することが遺伝子学的手法により確認された。</p> <p>齧歯類の重症疾患の原因となる picornavirus 科に属する cardiovirus は、その罹患率、多様性、ヒトでの症状等についてはあまり知られていない。発熱のある乳児の便検体から 1981 年に培養された Saffold virus は、cardiovirus に分類されている。今回、患者検体から直接ヒト cardiovirus をクローニングしたことについて報告する。これはインフルエンザ様の症状を示した子供の呼吸分泌物から pan-viral microarray 法を用いて発見した最初の報告である。ほぼ全長のウイルスゲノム(7961 bp)の系統樹解析で、ウイルスは cardiovirus のサブグループである Theiler's murine encephalomyelitis virus(TMEV) に属し、Saffold virus と最も密接に関係があった。719 の呼吸器サンプル(急性呼吸器症状を示した患者からは 637 検体(89%))と神経系疾患患者(無菌性髄膜炎、脳炎及び多発性硬化症)からの髄液検体 400 の RT-PCR によるスクリーニングでは、cardiovirus 感染の痕跡は認められなかった。しかし、胃腸炎患者 498 人の排泄物 751 検体のスクリーニングの結果、6 検体より cardiovirus (1.2%) が検出された。これら Saffold virus を含む 8 つのヒト cardiovirus は、系統樹解析によりすべて同じところにクラスターされたが、VP1 遺伝子にかなりの多様性が認められた(アミノ酸の相同性は 66.9%-100%)。これらの結果は、これまでほとんど確認されていなかったが、現在は主に消化管において確認され、無症候で排出され、そして腸内外の疾患に関連している可能性がある新しいヒト TMEV 様の cardiovirus の多様な集団が存在することを示唆している。</p>				使用上の注意記載状況・ その他参考事項等				
	<table border="1"> <tr> <td>報告企業の意見</td> <td>今後の対応</td> </tr> <tr> <td>別紙のとおり</td> <td>今後とも関連情報の収集に努め、本剤の安全性の確保を図っていきたい。</td> </tr> </table>					報告企業の意見	今後の対応	別紙のとおり	今後とも関連情報の収集に努め、本剤の安全性の確保を図っていきたい。
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101

一 般 的 名 称	①人血清アルブミン、②人血清アルブミン、③人血清アルブミン*、④人免疫グロブリン、⑤乾燥ペプシン処理人免疫グロブリン、⑥乾燥スルホ化人免疫グロブリン、⑦乾燥スルホ化人免疫グロブリン*、⑧乾燥濃縮人活性化プロテインC、⑨乾燥濃縮人血液凝固第Ⅷ因子、⑩乾燥濃縮人血液凝固第Ⅸ因子、⑪乾燥抗破傷風人免疫グロブリン、⑫抗HBs人免疫グロブリン、⑬トロンビン、⑭フィブリノゲン加第ⅩⅢ因子、⑮乾燥濃縮人アンチトロンビンⅢ、⑯ヒスタミン加入免疫グロブリン製剤、⑰人血清アルブミン*、⑱人血清アルブミン*、⑲乾燥ペプシン処理人免疫グロブリン*、⑳乾燥人血液凝固第Ⅸ因子複合体*、㉑乾燥濃縮人アンチトロンビンⅢ
販 売 名 ( 企 業 名 )	①献血アルブミン20“化血研”、②献血アルブミン25“化血研”、③人血清アルブミン“化血研”*、④“化血研”ガンマーグロブリン、⑤献血静注グロブリン“化血研”、⑥献血ベニコロン-I、⑦ベニコロン*、⑧注射用アナクトC2,500単位、⑨コンファクトF、⑩ノバクトM、⑪テタノセーラ、⑫ヘパトセーラ、⑬トロンビン“化血研”、⑭ボルヒール、⑮アンスロビンP、⑯ヒスタグロビン、⑰アルブミン20%化血研*、⑱アルブミン5%化血研*、⑲静注グロブリン*、⑳ノバクトF*、㉑アンスロビンP1500注射用
報 告 企 業 の 意 見	<p>cardiovirus は、picornavirus 科に分類される属名の一つである。cardiovirus 属のウイルスはエンベロープを持たず、直径約 30nm で正 20 面体のカプシッドを持ち、核酸は一本のプラス鎖 RNA である。cardiovirus 属には次の 2 つのサブグループがある；脳心筋炎ウイルス (encephalomyocarditis virus ; EMCV)、タイラーのマウス脳脊髄炎ウイルス (Theiler's murine encephalomyelitis virus ; TMEV)。これらのウイルスは、げっ歯類に感染し消化器官で増殖した後、糞便経口ルートで伝播する。ウイルスが腸管感染しても大抵は軽度か無症状であるが、腸管外に拡がると全身性の疾患を惹き起こす。EMCV 系統のウイルスは脳炎及び心筋炎を惹き起こし、TMEV 系統のウイルスは中枢神経系感染に関連している。ヒトから分離されたとされる cardiovirus 属のウイルスも報告されているが、ヒトから直接クローニングされたことはなく、その罹患率、多様性、ヒトでの症状等についてはあまり知られていない。</p> <p>本剤の製造工程には、冷エタノール分画工程、ウイルス除去膜ろ過工程あるいは加熱工程等の原理の異なるウイルス除去及び不活性化工程が存在しているので、ウイルスクリアランスが期待される。各製造工程のウイルス除去・不活性化効果は、「血漿分画製剤のウイルスに対する安全性確保に関するガイドライン (医薬発第 1047 号、平成 11 年 8 月 30 日)」に従い、ウシウイルス性下痢ウイルス (BVDV)、仮性狂犬病ウイルス (PRV)、ブタパルボウイルス (PPV)、A 型肝炎ウイルス (HAV) または脳心筋炎ウイルス (EMCV) をモデルウイルスとして、ウイルスプロセスバリデーションを実施し、評価を行っている。今回報告した cardiovirus 属には、モデルウイルスとして使用している EMCV そのものが属しており、上記バリデーションの結果から、本剤の製造工程が EMCV の除去・不活性化効果を有することを確認している。また、これまでに本剤による cardiovirus 感染の報告例は無い。</p> <p>以上の点から、本剤は cardiovirus に対する安全性を確保していると考える。</p>

\*現在製造を行っていない

# Identification of cardioviruses related to Theiler's murine encephalomyelitis virus in human infections

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Cardioviruses comprise a genus of picornaviruses that cause severe illnesses in rodents, but little is known about the prevalence, diversity, or spectrum of disease of such agents among humans. A single cardiovirus isolate, Saffold virus, was cultured in 1981 in stool from an infant with fever. Here, we describe the identification of a group of human cardioviruses that have been cloned directly from patient specimens, the first of which was detected using a pan-viral microarray in respiratory secretions from a child with influenza-like illness. Phylogenetic analysis of the nearly complete viral genome (7961 bp) revealed that this virus belongs to the Theiler's murine encephalomyelitis virus (TMEV) subgroup of cardioviruses and is most closely related to Saffold virus. Subsequent screening by RT-PCR of 719 additional respiratory specimens [637 (89%) from patients with acute respiratory illness] and 400 cerebrospinal fluid specimens from patients with neurological disease (aseptic meningitis, encephalitis, and multiple sclerosis) revealed no evidence of cardiovirus infection. However, screening of 751 stool specimens from 498 individuals in a gastroenteritis cohort resulted in the detection of 6 additional cardioviruses (1.2%). Although all 8 human cardioviruses (including Saffold virus) clustered together by phylogenetic analysis, significant sequence diversity was observed in the VP1 gene (66.9%–100% pairwise amino acid identities). These findings suggest that there exists a diverse group of novel human Theiler's murine encephalomyelitis virus-like cardioviruses that hitherto have gone largely undetected, are found primarily in the gastrointestinal tract, can be shed asymptotically, and have potential links to enteric and extraintestinal disease.

DNA microarrays | gastroenteritis | influenza-like illness | picornavirus | virus discovery

**P**icornaviruses are positive single-stranded RNA viruses that cause a variety of important disease states in humans and animals. Several genera of picornaviruses are recognized, based on genomic sequence and virus biology. The *Cardiovirus* genus of the family Picornaviridae consists of two subgroups: Theiler's murine encephalomyelitis virus (TMEV) and related viruses (Theiler-like virus NGS910 of rats, Vilyuisk virus) (1–3), and encephalomyocarditis virus (EMCV) and related viruses (EMCV, Mengovirus, Columbia SK virus, Maus–Elberfeld virus) (4). All these viruses infect rodents, replicate in the gastrointestinal (GI) tract and are transmitted by the fecal–oral route. Although enteric infection by these viruses is often mild or asymptomatic, extraintestinal spread of these viruses can occur and can lead to systemic disease (1). As their name implies, the EMCV-like agents cause encephalitis and myocarditis, whereas the TMEV family is linked to CNS infection. In experimental settings, intracerebral inoculation of mice with TMEV can produce acute encephalomyelitis and/or a chronic demyelinating disease resembling human multiple sclerosis (MS), depending upon the strain of TMEV used (5). Oral

inoculation with TMEV may also result in encephalomyelitis, especially when large inocula are delivered to neonatal mice (6).

Whether authentic human cardioviruses exist has long been debated. The first candidate human cardiovirus was Vilyuisk virus, which was linked to Vilyuisk encephalitis, an unusual neurodegenerative disease found among the Yakuts people of Siberia in the 1950s and still endemic to the region (7, 8). The Vilyuisk virus was initially isolated from the cerebrospinal fluid (CSF) of an affected patient and underwent 41 serial passages in mice before sequencing and characterization as a TMEV-like picornavirus (3, 9). Given its sequence similarity to TMEV and its extensive passage history in mice, questions have arisen as to whether the virus may in fact be of murine origin. In 1981, another TMEV-related cardiovirus was cultured from the stool of an infant who presented with a febrile illness (10). Although early passages appeared to show that the virus was transmissible, long-term continuous propagation of the isolate has been problematic. The nearly complete genomic sequence of this isolate (provisionally called Saffold virus) was recovered from frozen stocks by cloning in 2007 and was found to be much more divergent from TMEV than Vilyuisk virus (10). However, neither Vilyuisk nor Saffold virus was cloned directly from primary clinical specimens, and the diversity, prevalence, and potential clinical manifestations of human cardiovirus infection have remained largely unexplored.

We have previously developed a pan-viral DNA microarray (Virochip; University of California, San Francisco) designed to detect known and novel viruses in clinical specimens on the basis of homology to conserved regions of known viral sequences (11). The current study uses microarrays from the third and fourth generations of this platform (Viro3, Viro4). The Viro3 platform has 19,841 viral oligonucleotides derived from all publicly available viral sequence as of June 2004 (12, 13). The Viro4 platform is a streamlined update of the Viro3 platform consisting of 14,740 viral oligonucleotides derived from all publicly available viral sequence as of June 2006. The Virochip has been used to detect novel pathogens such as the severe acute respiratory syndrome coronavirus (14) and XMRV, a retrovirus identified in prostate tissue of men with germ-line mutations in RNase L (15). The platform has also been successfully used to detect

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known and divergent viruses in acute respiratory tract infections in several recently published studies (12, 13, 16, 17).

In this study, we used the Virochip to screen respiratory secretions from patients with influenza-like illness who lacked a diagnosis despite extensive microbiological testing. In one such patient, we detected and fully sequenced a cardiavirus in the Saffold group. Related cardiaviruses were subsequently found in stool specimens from an additional six individuals collected as part of a study examining household transmission of gastroenteritis (18). We report here the existence and overall phylogeny of a diverse group of human cardiaviruses and discuss their potential association with human disease.

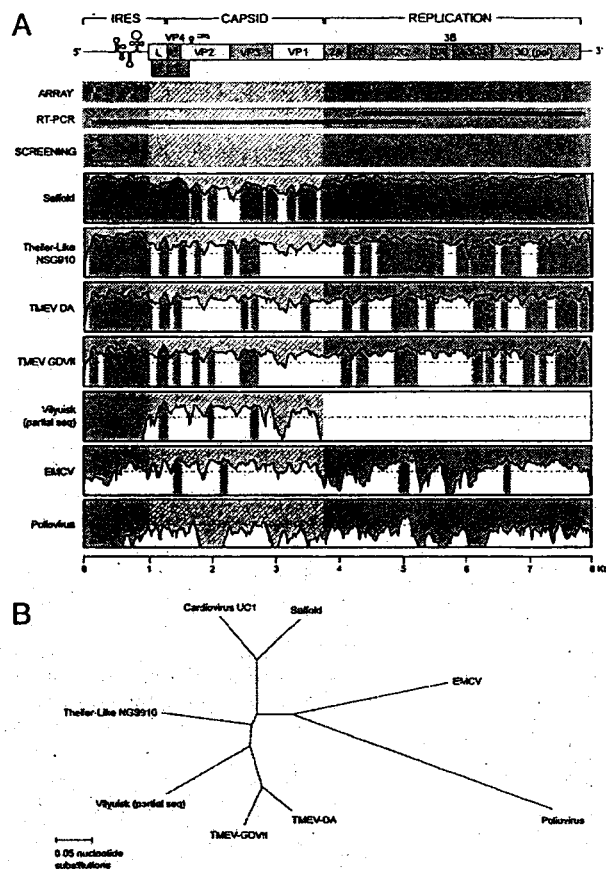
## Results

**Detection of a Cardiavirus in a Patient with Influenza-Like Illness.** A total of 460 respiratory secretions from patients meeting a case definition of influenza-like illness were screened for respiratory viruses by culture. In 108 culture-negative specimens selected from elderly and pediatric patients, 16 specimens remained negative after subsequent RT-PCR testing for respiratory syncytial virus (RSV), influenza A/B (Flu A/B), rhinovirus (RV), and enterovirus (EV). These 16 specimens were assayed for the presence of viruses using the Virochip (Viro3), with microarray analysis carried out using E-Predict and ranked z score analysis, as previously described (12, 19).

Four of the 16 specimens yielded a positive microarray hybridization signature suggestive of a virus. Two of the signatures corresponded to metapneumovirus, one signature corresponded to adenovirus, and one signature indicated the presence of a cardiavirus related to TMEV. From the microarray containing the cardiavirus signature, the highest intensity oligonucleotides mapped to the 5'-untranslated region (5'-UTR) and 2C gene of the TMEV genome, the most conserved regions among cardiaviruses and picornaviruses in general (Fig. 1A, "ARRAY"). To recover viral sequence, we designed primers based on the highest intensity array features and alignment of well conserved sequences from four cardiaviruses (TMEV-DA, TMEV-GDVII, Theiler-like NGS910 virus, and EMCV). One set of primers successfully amplified a 224-bp fragment from the viral 5'-UTR. The fragment shared 90% nucleotide identity with the 5'-UTR region of Theiler-like NGS910 virus. This finding established that the virus in question was indeed a cardiavirus and a relative of the TMEV group of viruses. We designated this initial cardiavirus strain UC1.

**Complete Genome Sequencing and Analysis of UC1.** To clone and sequence the remainder of the UC1 genome, additional short fragments were first obtained from conserved regions in the 2C (helicase) and 3D (polymerase) genes by use of consensus PCR primers derived from alignment of the four cardiaviruses mentioned previously. Long-range RT-PCR using specific primers was then used to bridge the gaps. This resulted in PCR amplification of two long overlapping fragments (~5.3 and 3.7 kB in size) jointly spanning nearly the entire length of the virus genome (Fig. 1A, "RT-PCR"). Cloned ends of the genome were recovered and sequenced using a RACE amplification protocol (20, 21).

The nearly complete sequence of UC1 is 7961 nt in length and forms a distinct branch in the *Cardiavirus* genus with Saffold virus (Fig. 1B). The overall nucleotide identity to Saffold virus is >90% in the 5'-UTR and the region coding for the nonstructural proteins but only 70% in the region coding for the capsid proteins (Fig. 1A, "Saffold"). There is much less overall nucleotide sequence identity to other members of the TMEV subgroup (70–80%) and EMCV (50–55%). A poly(C) tract that has been reported in EMCV but not in TMEV strains is not present in the 5'-UTR of UC1. Similar to other cardiaviruses, the ORF of UC1 is predicted to code for a single 2296-amino acid polyprotein that is subsequently cleaved into the L protein, the



**Fig. 1.** Genome sequence of UC1. (A) Genome sequence similarity plots compare UC1 with Saffold virus, Theiler-Like NGS910 virus, TMEV-DA, TMEV-GDVII, Vilyuisk virus (partial sequence only), EMCV, and poliovirus. The y axis scale for each plot represents percentage of nucleotide identities from 0% to 100%. Regions of the genome with percentage of nucleotide identities of >70% are highlighted in pink. The Virochip oligonucleotides used to detect UC1 ("ARRAY"), the fragments generated by long-range RT-PCR and used to sequence most of the virus ("RT-PCR"), and the cardiavirus primers and resulting PCR fragments used for screening of stool, CSF, and respiratory secretions ("SCREENING") are also shown mapped onto the UC1 genome. The sequences of these primers are provided in Table S1. (B) Radial tree depicts the phylogenetic relationships between the genomes of UC1 and the seven aforementioned cardiaviruses.

capsid proteins (VP1, VP2, VP3, and VP4), and nonstructural proteins involved in viral replication (2A, 2B, 2C, 3A, 3B, 3C, and 3D) (Fig. 1A). Like Saffold virus, UC1 encodes an L protein containing a zinc finger, an acidic domain, and a partially deleted Ser/Thr-rich domain (22, 23) and potentially encodes a severely truncated L\* protein that begins with an ACG codon rather than AUG (22, 23) [supporting information (SI) Fig. S1A].

In cardiaviruses, the surface loops CD of VP1 and EF of VP2 are exposed on the capsid surface and are thought to be involved in host cell tropism and viral pathogenesis (24). These loops are the regions of greatest divergence between UC1 and the other cardiaviruses, including Saffold virus (Fig. S1B). Between UC1 and Saffold virus, there is 52% and 61% amino acid identity in the exposed surface loops CD and EF, respectively. The corresponding identities (29% and 24%) are much lower between UC1 and the rodent cardiaviruses.

**Comparison of UC1 Amino Acid Sequence with Other Cardiaviruses.** The level of divergence between the sequence of UC1 and other cardiaviruses is maintained at the amino acid level. Between UC1

Table 1. Amino acid identity of predicted UC1 proteins

Gene	Predicted size, aa	Percent amino acid identity to					
		Saffold	NGS910	Vilyuisk	TMEV-DA	EMCV	Polio
L protein	71	77	61	60	60	26	0
VP4	72	99	72	72	68	62	19
VP2	269	83	69	67	71	64	30
VP3	231	85	80	76	75	68	28
VP1	275	77	56	55	59	48	14
Nonstructural	1389	98	91		83	40	22
Polyprotein	2296	91	76		71	52	22

and Saffold virus, the capsid proteins VP1, VP2, and VP3 are only 77–85% identical, whereas the nonstructural proteins are highly conserved (98% overall identity) (Table 1). The amino acid identities between UC1 and its closest rodent relatives (NGS910 virus and TMEV) are much lower, 56–80% for the capsid proteins and 83–91% for the nonstructural proteins. These comparisons confirm that UC1 is most closely related to Saffold virus, although there is significant sequence divergence in the capsid proteins containing the putative receptor binding sites.

**Prevalence of Cardioviruses in Clinical Specimens.** To investigate the prevalence of cardiavirus infection in acute human illnesses, we designed PCR primers targeting the 5'-UTR to amplify cardiaviruses by real-time one-step RT-PCR. In our initial screen, we ran two RT-PCRs using conserved primers designed to amplify 102-bp and 224-bp fragments from the 5'-UTR of UC1, Saffold virus, or all mouse strains of TMEV. By probit analysis (i.e., the concentration of the target sequence testing positive in 95% of cases) using *in vitro* transcribed UC1 mRNA, the sensitivity of the RT-PCR assay for detection of cardiaviruses was 600 copies. Standard curves generated using pooled cardiavirus-negative specimens spiked with UC1 mRNA were linear from 10<sup>4</sup> to 10<sup>11</sup> copies/ml ( $R^2 = 0.9831-0.9944$ , Fig. S2). The presence of PCR inhibitors was estimated to be <3% by yeast RNA spiking experiments on randomly selected stool specimens (only 2 of 95 RT-PCRs failed to amplify the yeast positive control). All positives in the initial screen were sequenced and then further confirmed by another RT-PCR using primers designed to amplify an overlapping 608-bp fragment (Fig. 1A, "SCREENING").

Since UC1 was first identified in respiratory secretions, we screened 719 respiratory specimens from two large groups of patients: 278 nasopharyngeal aspirates from pediatric patients at a single hospital (190 specimens from patients with an acute respiratory illness) (13) and 441 pooled oropharyngeal and nasopharyngeal swabs from individuals in California with influ-

enza-like illness (25). None of the 719 total respiratory specimens tested was positive for cardiaviruses.

We next conducted screening of CSF specimens from patients with aseptic meningitis ( $n = 60$ ), patients with encephalitis ( $n = 300$ ), and patients with MS ( $n = 40$ ) for cardiaviruses by RT-PCR. None of the 400 CSF specimens tested was found to be positive.

Given the prominent association of picornaviruses with enteric infection and the known fecal-oral route of transmission, we then sought to assess the prevalence of human cardiaviruses in stool. We examined 751 stool specimens from 498 individuals collected as part of a cohort study of household transmission of *Helicobacter pylori* and gastroenteritis (18). The vast majority of subjects were children, with 443 (89%) children younger than 5 years, 30 (6%) children between 5 and 18 years, and 25 (5%) adults. Specimens from 6 children (1.2% of the 498 individuals) were positive for cardiaviruses (strains UC2–UC7). All cardiavirus-positive stool specimens were from children <2 years old and from different households. Symptoms in the 6 children included diarrhea and vomiting in 3 (50%) and diarrhea only in 1 (17%); the remaining 2 children were asymptomatic. Of note, from 2 of the symptomatic children, one with diarrhea and vomiting and the other with diarrhea, a cardiavirus was identified not during acute illness but in a specimen obtained months after each child had recovered.

To investigate the possibility of coinfection with additional viruses, we used the Virochip (Viro4) to analyze the nine available specimens collected from the six cardiavirus-positive cases (Table 2). As expected, all six cardiavirus-positive cases were positive for a cardiavirus by Virochip. In three of the cardiavirus-positive stool specimens, there was evidence of coinfection: in two specimens by caliciviruses (norovirus and sapovirus) and in one specimen by a rotavirus. In the other three individuals, viruses other than cardiaviruses were detected in the stool at the time of the first visit (adenovirus, norovirus and parechovirus, norovirus and enterovirus), but only cardiavirus

Table 2. Patients with stool positive for cardiaviruses

ID	Age at first visit, months	Number ill in household	Days between visits 1 and 2	Clinical symptoms		Virochip/PCR results	
				10 days prior to visit 1	Between visits 1 and 2	Visit 1	Visit 2
UC2	8.4	4/10	—	Diarrhea/vomiting	—	Cardiavirus, rotavirus	—
UC3	6.1	1/5	139	None	none	—	Cardiavirus, norovirus
UC4	21.4	1/9	91	None	none	Adenovirus	Cardiavirus
UC5	16.3	6/6	95	Diarrhea/vomiting	none	Norovirus, parechovirus	Cardiavirus
UC6	14.0	1/5	—	Diarrhea/vomiting	—	Cardiavirus, sapovirus	—
UC7	18.6	3/7	94	Diarrhea	none	Norovirus, enterovirus	Cardiavirus

Dashes indicate entries for which data and/or specimens were not available.

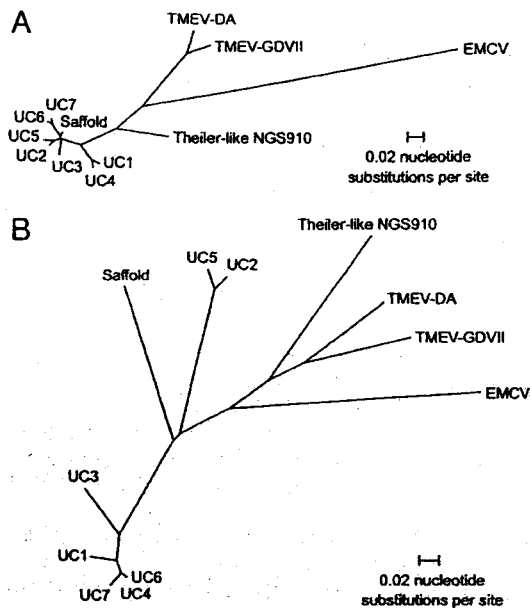


Fig. 2. Strain variation of human cardiomyoviruses. (A) Radial tree of a 608-bp region within the 5'-UTR. (B) Radial tree of an 819-bp region corresponding to the VP1 gene. Strain designations UC2 to UC7 correspond to patients as listed in Table 2.

was detected in the stool by the second visit. All Virochip results were subsequently confirmed by PCR and sequencing using virus-specific primers.

To assess the sequence variation within different cardiomyovirus strains, we analyzed a 608-bp region from the 5'-UTR and an 819-bp region corresponding to the VP1 gene for the six positive cardiomyovirus cases (Fig. 2). The sequence variations within the 608-bp region from the 5'-UTR (2.0–9.1%) and within the 819-bp region corresponding to the VP1 gene (0.3–36.7%) were consistent with infection by independently acquired cardiomyovirus strains. The amino acid sequence identities in the VP1 gene were lowest between UC2/UC5 and the other cardiomyoviruses (66.9% for Saffold virus, 71.0–72.8% for the other UC strains).

### Discussion

Using a pan-viral microarray, we analyzed 16 respiratory specimens from patients with influenza-like illness who still lacked a diagnosis after extensive tests for respiratory viruses. In one specimen, we found a signature for a cardiomyovirus. Sequence recovery of the genome and phylogenetic analysis revealed that this virus (UC1) is divergent from the rodent cardiomyoviruses and clusters with the Saffold agent. Like Saffold virus, UC1 may code for a truncated L\* protein (Fig. S1A) that has been implicated in viral persistence and chronic infection of the CNS in TMEV (26). However, because the L\* protein of UC1 begins with an ACG codon rather than AUG, it is unclear whether any functional protein is actually expressed, although small amounts of L\* protein have been detected in TMEV strains carrying the ACG codon (27).

The binding of sialic acid to TMEV is strongly associated with persistence and neurovirulence, and three amino acids in the VP2 protein are directly involved in this interaction (28, 29). In both UC1 and Saffold virus, there is a substitution or deletion at each of these three positions (Fig. S1B), suggesting that sialic acid is unlikely to serve as a receptor for these viruses. Although the cellular receptor is presently unknown, the sequences of UC1 and Saffold virus are most divergent in the capsid region, sharing only 77% and 83% amino acid identity in the VP1 and VP2 proteins, respectively, and 52%

and 61% identity in the exposed surface loops CD and EF, respectively. These differences may reflect the use of distinct cellular receptors or may be the result of immune selection during virus evolution (or both); further studies will be required to shed light on these issues.

Cardiomyoviruses were detected in six children out of a total of 498 individuals (1.2%) enrolled in a large gastroenteritis study. Although the initial specimen that was used to culture Saffold virus was collected >25 years ago, cardiomyoviruses UC1 through UC7 were collected from 2000 to 2006, indicating that human cardiomyoviruses continue to circulate in the population. Despite the use of screening RT-PCR assays able to detect all strains of TMEV, cardiomyoviruses detected in human clinical specimens clustered together and were phylogenetically distinct from the rodent cardiomyoviruses (Fig. 2).

Further studies will be required to define the pathogenic role of cardiomyovirus infection in the intestine fully. Although we did recover a cardiomyovirus from a number of cases with symptomatic enteritis, other potential GI pathogens were also detected in these cases. Thus, it is presently unclear how frequently enteric cardiomyoviral infection produces clinical illness. Moreover, we detected cardiomyoviruses in stool from subjects without enteritis, suggesting that asymptomatic shedding of cardiomyoviruses in the GI tract can and does occur. In this respect, cardiomyovirus infection in humans may mimic that of murine TMEV, which is often shed asymptotically in naturally acquired infections (30).

Cardiomyovirus infection outside the GI tract is sometimes associated with severe disease in rodents, including encephalomyelitis, demyelinating disease, and myocarditis (1), although only a small percentage of mice naturally infected with TMEV develop systemic disease (1, 5). Our wider screening for cardiomyoviruses indicates that cardiomyovirus infection is uncommon in the setting of acute respiratory or neurological disease (e.g., aseptic meningitis, encephalitis, MS). However, while this manuscript was under review, Abed and Boivin (31) reported detection of Saffold-like cardiomyoviruses in three children with acute respiratory illness. Moreover, in a case of influenza-like illness reported here, a cardiomyovirus was the sole agent identified despite comprehensive testing with culture, PCR, and a pan-viral microarray, suggesting that cardiomyoviruses may be pathogenic outside the GI tract in at least some instances.

One remarkable finding from this study was the diversity of the human cardiomyoviruses that have been identified. For the family Picornaviridae, the definition of a new species in a genus is having <70% amino acid identity in the coding regions of either VP1, 2C, 3C, or 3D (32). By this strict definition, cardiomyoviruses UC2 and UC5 would classify as a novel species distinct from Saffold virus, with 66.9% amino acid identity in the VP1 gene. However, since cardiomyoviruses UC1 through UC7 and Saffold virus as a whole clearly define a separate group within the *Cardiomyovirus* genus by phylogenetic analysis (Figs. 1B and 2), we propose a systematic nomenclature for the human cardiomyoviruses, designating all members of this group HTCv, for human TMEV-like cardiomyovirus, and referring to the strains in this group by a brief suffix (e.g., Saffold agent would be designated HTCv-Saf, UC1 would be designated HTCv-UC1).

Several lines of evidence support the inference that HTCvs are bona fide human viruses and not the products of sporadic viral cross-over events from rodents to humans: (i) all seven cardiomyoviruses from humans in this study are strains of HTCv, with no mouse TMEV sequences detected in 1870 total clinical specimens despite screening using two consensus PCR primer sets designed to amplify UC1, Saffold virus, or mouse TMEV; (ii) sequence variations within HTCv UC1–7 are most consistent with independent acquisition of different virus strains by patients; and (iii) HTCv is substantially diverged from the rodent cardiomyoviruses, especially in the capsid region containing

the putative receptor binding sites. Taken together, our findings indicate that HTCVs are novel human picornaviruses in the *Cardiovirus* genus that are found primarily in the GI tract, can be shed asymptotically, and have potential links to self-limited enteric disease and, rarely, to influenza-like illness. Although the full spectrum of clinical diseases linked to HTCv and the mechanisms underlying viral replication remain to be elucidated, the studies reported here now open all these questions to direct experimental scrutiny.

## Materials and Methods

**Clinical Specimens.** *Respiratory secretions from the California Influenza Surveillance Program study.* A total of 943 respiratory specimens were sent to the California Department of Health Services (DHS) during the 2005–2006 season (25). Among these 943 specimens, 460 were pooled nasopharyngeal and oropharyngeal swabs collected as part of the California Influenza Surveillance Program (CISP) study under protocols approved by the DHS. Patients enrolled in the CISP study fulfilled a clinical case definition of influenza-like illness (temperature of 37.8°C or greater and a cough and/or sore throat in the absence of a known cause other than influenza). Sixty percent, or 280 specimens, were positive for a virus by culture. Among the remaining 180 culture-negative specimens, a subset of 108 specimens selected from elderly and pediatric patients was then subjected to further screening by RT-PCR to exclude cases of RSV, Flu A/B, RV, and EV (33). Sixteen specimens negative by culture and RT-PCR were then examined using the Virochip. We subsequently screened 441 CISP specimens with remaining available specimen material (96% of the 460 total collected specimens) for cardioviruses by RT-PCR.

*Respiratory secretions from the UCSF pediatric respiratory infections study.* This collection consisted of 278 consecutive nasopharyngeal aspirates from pediatric patients seen at UCSF from December 2003 to June 2004 (13). All specimens were collected under protocols approved by the UCSF Institutional Review Board. In this group, 190 of the patients (68%) had a respiratory illness, defined as an upper respiratory infection, bronchiolitis, croup, asthma exacerbation, or pneumonia. The remaining 88 patients (32%) were asymptomatic.

*Stool from the Stanford Infection and Family Transmission cohort.* The Stanford Infection and Family Transmission (SIFT) cohort of 4333 individuals was initiated in 1999 to evaluate the association between *H. pylori* infection and gastroenteritis transmission prospectively (18). Among the 3063 subjects who consented to further use of biological specimens, 774 stool specimens were obtained from 514 individuals; of those, 751 specimens from 498 subjects were available for study. Additional details on the 751 specimens screened for cardioviruses by RT-PCR are described in *SI Text*.

*CSF specimens from patients with aseptic meningitis, encephalitis, and MS.* A total of 60 CSF specimens from patients with clinically diagnosed aseptic meningitis, 300 CSF specimens from patients with encephalitis (who lacked a diagnosis despite comprehensive testing) (34), and 40 CSF specimens from patients with MS were screened for cardioviruses by RT-PCR. Specimens were collected under protocols approved by the California DHS (encephalitis specimens) or the UCSF Institutional Review Board (aseptic meningitis and MS specimens).

**Specimen Preparation and Diagnostic Testing.** In the CISP study, routine tube culture or shell vial culture of pooled nasopharyngeal and oropharyngeal swab specimens followed by specific monoclonal antibody testing for viral identification was performed as previously described (33, 35). Total nucleic acid was then extracted from the specimens using the MasterPure Complete DNA and RNA Purification Kit (Epicentre). Real-time one-step RT-PCR assays for RSV, Flu A/B, and picornavirus (inclusive of RV and EV) were then performed as previously described (25, 33, 36). In the UCSF pediatric respiratory infections study, 200- $\mu$ l aliquots of nasopharyngeal lavage were used to extract RNA using the RNeasy Mini Kit (Qiagen Corporation), including on-column DNase digestion. In the SIFT cohort, stool was suspended in 2 ml of PBS at 10% weight per volume and the PureLink96 Viral RNA/DNA Kit (Invitrogen) was used to extract RNA for RT-PCR and Virochip analysis. Cerebrospinal fluid specimens were processed using either a Zymo MiniRNA Isolation Kit (Zymo Research) or the MasterPure Complete DNA and RNA Purification Kit.

Virochip analysis of CISP and SIFT specimens was carried out as previously described (14). Extracted nucleic acid specimens were amplified and labeled using a Round A/B protocol and were hybridized to the Virochip. Microarrays (National Center for Biotechnology Information GEO platforms GPL3429 for Viro3 and GPL6862 for Viro4) were scanned with an Axon 4000B scanner (Axon Instruments). Virochip results were analyzed using cluster analysis, E-Predict,

and z score analysis as previously described (12, 19, 37). All Virochip microarrays have been submitted to the GEO database (National Center for Biotechnology Information GEO series number GSE11569, accession numbers GSM291246–GSM291254).

**Complete Genome Cloning and Sequencing (UC1 strain).** Conserved primers from the 5'UTR of cardioviruses were designed based on the highest intensity microarray oligonucleotides and alignment of well conserved sequences from four cardioviruses for which full genome sequences were available: TMEV-DA, TMEV-GDVII, Theiler-like NGS910 virus, and EMCV. After short viral fragments were obtained, six sets of specific primers derived from sequenced fragments and conserved primers were then used to sequence the genome by long-range RT-PCR and 5'/3' RACE (rapid amplification of cDNA ends). Amplicons for sequencing were cloned into plasmid vectors using the TOPO TA Cloning System (Invitrogen) and sequenced on an ABI3130 Genetic Analyzer (Applied Biosystems) using standard Big Dye terminator (version 3.1) sequencing chemistry. The completed genome sequence of UC1 has been deposited into GenBank (GenBank accession number EU376394).

**Phylogenetic Analysis (UC1 strain).** Nucleotide and protein sequences associated with the following reference virus genomes were obtained from GenBank: Saffold virus (NC\_009448), TMEV-DA (M20301), TMEV-GDVII (NC\_001366), Theiler-like NGS910 virus (AB090161), EMCV (NC\_001479), poliovirus (NC\_002048), and the partially sequenced genome of Vilyuisk virus (M94868). For amino acid analysis, ORFs predicted using ORF Finder (National Center for Biotechnology Information) were used. Multiple sequence alignment was performed using ClustalX (version 1.83). Neighbor-joining trees using the Kimura two-parameter distance correction were generated using 1000 bootstrap replicates and displayed using MEGA (version 3.1). Sequence identities were calculated using BioEdit (version 7.0.9.0).

**RT-PCR Screening for Cardioviruses.** Real-time quantitative RT-PCR (qRT-PCR) screening for cardioviruses with SYBR Green I (Invitrogen) was performed using conserved PCR primer sets CardioUTR-1F/CardioUTR-2R-A and CardioUTR-1F/CardioUTR-2R-B (Table S1) on a DNA Engine Opticon System (Bio-Rad). To determine limits of sensitivity of the qRT-PCR assay, probit analysis of results from 10 qRT-PCR replicates of eight serial half-log dilutions of *in vitro* transcribed UC1 mRNA (from a starting concentration of  $\sim 10^5$  copies/ml) was performed using StatsDirect (StatsDirect Ltd.). Standard curves of the qRT-PCR assay were calculated from 3 qRT-PCR replicates of seven serial log dilutions of RNA extracted from pooled respiratory secretions, stool suspensions, and PBS spiked with UC1 RNA (10 specimens per pool). To assess for the presence of PCR inhibitors, RT-PCR for yeast was carried out on 95 randomly selected stool samples, each spiked with 1 ng of *in vitro* transcribed *Saccharomyces cerevisiae* intergenic RNA as a positive control (38).

Positive bands corresponding to the expected 102-bp and 224-bp amplicons were cloned and sequenced in both directions using vector primers M13F and M13R. Secondary confirmation of all positive reactions was performed using RT-PCR with primers CardioUTR-1F and CardioUTR-3R (Table S1), which generated a larger 608-bp amplicon, also in the 5'-UTR. To obtain the full sequences of the VP1 gene in strains UC2 through UC7, RT-PCRs were performed using conserved primers flanking the VP1 region of UC1 and Saffold virus (Table S1). The sequences of the 5'-UTR and VP1 amplicons corresponding to cardiovirus strains UC2 through UC7 have been deposited in GenBank (accession numbers EU604739–EU604750).

**PCR Confirmation for Virochip-Positive Stool Specimens.** All nine specimens collected from the six positive cardiovirus cases were analyzed using the Virochip as previously described (11, 12). Confirmatory PCR for calicivirus, adenovirus, and parechovirus was carried out using conserved primers as previously reported (39–41). Amplified PCR bands of the expected size were gel extracted and sequenced using standard BigDye chemistry on an ABI3130 (Applied Biosystems).

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研究報告の概要 109	<p>○ヘンドラウイルス感染、ヒト、ウマ-オーストラリア オーストラリア、ブリスベーンの動物病院スタッフがヘンドラウイルスに感染した。看護師1名と獣医師1名が、感染したウマを治療した後でウイルスに感染したと診断された。患者は2名とも重症である。潜伏期間は最大14日間で、スタッフの間から新たな患者が出ないか監視が続けられている。ウマは感染した3頭中1頭が死亡、1頭を安楽死させたが、1頭は回復に向かっている。ヘンドラウイルスが以前に流行したのは1994年で、ウマ14頭と調教師1名が死亡した。ヒト-ヒト感染の証拠はなく、広範囲に流行する危険性はない。</p>					<p>使用上の注意記載状況・ その他参考事項等</p> <p>合成血-LR「日赤」 照射合成血-LR「日赤」</p> <p>血液を介するウイルス、 細菌、原虫等の感染 vCJD等の伝播のリスク</p>
	<p>報告企業の意見</p> <p>オーストラリア、ブリスベーンの動物病院スタッフがヘンドラウイルスに感染し、重症となったとの報告である。ヘンドラウイルスはニパウイルスに近縁のウイルスで、現在のところオーストラリア以外での発生はない。</p>	<p>今後の対応</p> <p>日本赤十字社では、輸血感染症対策として問診時に海外渡航歴の有無を確認し、帰国(入国)後4週間は献血不適としている。今後も引き続き、新たなウイルス等による感染症の発生状況等に関する情報の収集に努める。</p>				



