Epidemiology of Human Adenoviruses: A 20-Year Retrospective Observational Study in Hospitalized Patients in Bern, Switzerland

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Background: Human adenovirus (HAdV) is an important pathogen seen in clinical practice. Long-term studies may help better understand epidemiological trends and changes in circulating genotypes over time.

Purpose: Using a large biobank of samples from hospitalized, adenovirus-positive patients over a 20-year period, we aimed to analyze long-term epidemiological trends and genotypic relatedness among circulating HAdV strains.

Methods: Based on samples from hospitalized patients confirmed to be HAdV positive in Bern, Switzerland, from 1998 to 2017, and on their associated demographic and clinical data, we identified epidemiological trends and risk factors associated with HAdV infection. HAdV genotyping was performed by PCR amplification and sequencing of the hypervariable hexon gene. The obtained sequences were phylogenetically compared with sequences from international HAdV strains.

Results: HAdV was identified in 1302 samples tested. Cases of HAdV infection were reported throughout the years with no clear seasonality. Upper respiratory tract samples, conjunctivitis swabs, and stool had the highest positivity rate (56.2%, 18.7%, and 14.2% of the cases, respectively). HAdV infection was highest among children ≤ 4 years old. Increased number of HAdV cases were observed in years 2009 (n = 110) and 2010 (n = 112). HAdV8 was the predominant genotype among patients older than 20 years, and was mostly associated with ophthalmic infection. Predominant genotypes among children ≤ 4 years old were HAdV1, HAdV2, and HAdV3, which were mostly associated with respiratory tract infections. Recurring peaks of increased HAdV cases were evidenced every 4 years among children ≤ 4 years old.

Conclusion: Our study gives novel insights on long-term epidemiological trends and phylogenetic relatedness among circulating HAdV strains in Switzerland, country in which little data on HAdV prevalence and diversity was so far available.

Keywords: adenoviruses, human, molecular epidemiology, clinical infections, genotype

Introduction

Human adenovirus (HAdV) is one of the most important viruses seen in clinical and public health today. This DNA virus is associated with a wide range of illnesses ranging from benign colds to more serious conditions including gastroenteritis, acute respiratory infections, conjunctivitis, haemorrhagic cystitis, meningencephalitis,1,2 and can cause severe disease in both immunocompetent3 and immunocompromised hosts, infections that sometimes result in death.2,4–7 HAdV is estimated to account for about 2% to 5% of all respiratory infections worldwide. This virus often causes epidemics and outbreaks,
especially when associated with either ophthalmic infections or respiratory tract infections.\textsuperscript{8-10} It is also known to be causative of emerging viral respiratory tract infections in humans,\textsuperscript{8,11-14} along with Middle East respiratory syndrome coronavirus, H7N9 influenza A virus or swine-like influenza H3N2 variant virus.

HAdV comprises over 100 genotypes recognized by the Human Adenovirus Working Group, July 2019, Update (http://hadvwg.gmu.edu) classified into seven species (HAdV-A to HAdV-G). In clinical diagnostics laboratories, molecular typing of HAdV positive samples is not routinely performed, unless epidemiological analyses are conducted. PCR-based techniques targeting genes encoding for the hypervariable hexon, fiber, penton base or DNA polymerase, and/or sequencing of their PCR products, allow identification of HAdV species and genotypes.

The recent emergence of the HAdV14 strains in the USA\textsuperscript{15} and Europe (Ireland)\textsuperscript{4} calls for a need to fill major gaps in understanding the molecular epidemiology and disease patterns of HAdVs. The predominant HAdV genotypes detected in association with clinical presentation differ among different countries and may change over time.\textsuperscript{16} Currently in Switzerland, the knowledge on the molecular epidemiology and clinical presentation of HAdV infection is lacking. This study presents the molecular epidemiology of HAdV and associated clinical findings based on a retrospective observational study from laboratory confirmed HAdV cases collected between 1998 and 2017, at the Institute for Infectious Disease (IFIK), following referral of specimens by clinicians from the largest hospitals and other regional clinics in Bern. Since outbreaks of HAdV have been reported in various countries,\textsuperscript{10,15,17-19} we therefore hypothesized that a thorough epidemiological analysis of the Bern HAdV clinical bio-bank, including genotyping and comparison to strains involved in international outbreaks, would provide information on potential outbreaks of HAdV that occurred in the last 20 years in Switzerland and how the epidemiological situation compares to that seen elsewhere in Europe during the studied period. To our knowledge, this is the first study to report HAdV epidemiology from a large clinical dataset over a 20-year period from patients of any age with positive HAdV infection. Overall, the study sheds light on epidemiological trends, risk factors for infection, and molecular epidemiology of HAdV circulating genotypes.

Materials and Methods

Laboratory Setting, Study Design and Ethical Approval

The institute for infectious diseases (IFIK), university of Bern, is a diagnostic, teaching and research institute in Switzerland. The institute covers the entire spectrum of microbiological diagnostics and also provides university education in clinical microbiology to students studying human medicine, dentistry, pharmacy, and cell biology. Samples used in this study are part of the IFIK clinical bio-bank and consisted of previously screened patient samples referred to the IFIK for diagnostics purposes by physicians from the Inselspital and other regional clinics in Bern. The clinical bio-bank consisted of patient samples and information on HAdV cases since 1998. The results of all cases were extracted from the electronic health-care records and information such as laboratory I.D., patient I.D., demographic data (age and sex), screening tests done, and sample type, which were available for most samples. Archived patients’ samples were approved for this research by the Swiss Ethics Committees on Research involving humans (BASEC-Nr: Req-2018-00158).

Study Population

HAdV positive infection was identified as sample(s) screened by either viral culture, immuno-chromatographic rapid test, immunofluorescence (IF), PCR (quantitative or qualitative real-time PCR), or all of the mentioned methods tested positive for HAdV by the department of clinical microbiology at IFIK. In instances in which more than one sample type was obtained from the same patient in a given year, multiple patient entries were removed and only the first screened positive sample type was kept for a given year. Similarly, subsequent HAdV infections from the same patient within a given year were excluded from epidemiological analyses.

Clinical Variables and Definitions

Patient age was defined at the date of sample collection. Clinical information and severity of HAdV positive cases were categorized as respiratory tract infection, fever only, ophthalmic infection, gastrointestinal infection, immunosuppressed, severe infection, co-infection, urinary tract infection, oncological disease, other manifestations, and unknown/no clinical information provided (Table 1).
Statistical Analysis
The Chi-square test was used to test for significant differences between categorical variables (test of independence). Logistic regression was used to determine significant years associated with increased HAdV cases. Data analysis was performed with the statistical R environment, version 1.0.153.

DNA Extraction, PCR, and Sequencing
Viral DNA was extracted from 200 µL of either original sample or cell culture supernatant of HAdV positive samples with the Boom extraction method20 or the NUCLISENS easyMAG (bioMérieux, Geneva, Switzerland) extractor, as per manufacturers instructions, and finally eluted in 50 µL. A PCR procedure that targeted the hypervariable regions 1–7 of the hexon gene was performed, according to Sarantis and colleagues.21 M13 universal priming tails (forward, 5’-TGTAAAACGACGGCCAGT-3’; and reverse, 5’-CAGGAAAACAGCTATGACCC-3’) were added to the Sarantis and colleagues primers (forward, 5’-CTGATGTAATGATACGGGCAAACAAGCAGCTATGACCC-3’; and reverse 5’-GGTTCGACGAAACCTATGACCC-3’). A standard protocol of the ZymoTaq Hotstart DNA PCR was applied, consisting of 2× Zymo reaction buffer, 0.2 mM dNTP mix, 0.5 µM of each primer, 5 U/µL ZymoTaq DNA Polymerase, 10 µL of DNA template and water up to 50 µL. Thermal cycling conditions were 95°C for 10 mins, 37 cycles at 95°C for 30 secs, 51°C for 30 secs, and 72°C for 1 min, with a final extension at 72°C for 7 mins. PCR products were resolved and visualized by electrophoresis on 1% agarose gel stained with RedSafe DNA Gel Stain (Invitrogen, Switzerland). Expected amplicon sizes ranged from 602 to 630 bp. Samples with PCR product band of the expected size were purified using 1.8× Agencourt Ampure XP beads (Beckman Coulter, Nyon, Switzerland), quantified by Qubit 3.0 fluorometer (ThermoFisher Scientific, Reinach, Switzerland), diluted to concentration of 180 ng in total reaction volume of 12 µL and then submitted to Microsynth AG (Balgach, Switzerland) for Sanger sequencing. All sequence data were deposited to the European Nucleotide Archive, under project reference PRJEB36200.

Sequence and Phylogenetic Analysis
The forward and reverse sequences were trimmed and assembled using DNASTAR Lasergene SeqMan Pro version: 7.1.0 (44.1). The consensus sequence of each sample obtained was blasted against all HAdV sequences available in GenBank to identify the species and genotype. The phylogenetic trees were constructed using the Neighbor-Joining method based on evolutionary distances computed using the number of differences method. MEGA-X22 was used to generate phylogenetic trees and evaluate topologies with 1000 bootstrap replicates.

Table 1 Definition of Clinical Categories Used in the Study

| Clinical Category | Clinical Features |
|-------------------|-------------------|
| Co-infection      | Respiratory syncytial virus, *Haemophilus influenzae* |
| Fever only        | Fever             |
| Gastrointestinal infection | Diarrhoea, vomiting, abdominal pain. Rare complications include haemorrhagic colitis, enteritis, hepatitis, cholecystitis, and pancreatitis |
| Immunosuppressed | Acute myeloid leukaemia, acute lymphatic leukaemia, chronic lymphatic leukaemia, esophagitis HIV stage III, after kidney transplant, after liver transplant, HIV positive, chemotherapy, immunosuppressed |
| Ophthalmic infection | Epidemic keratoconjunctivitis, pharyngoconjunctival fever, and nonspecific conjunctivitis |
| Other infection   | Exanthena, mononucleosis-like syndromes, leukocytosis, anaemia |
| Respiratory tract infection | Fever, pharyngitis, tonsillitis, cough, sore throat, pneumonia, bronchitis, rhinitis, angina, sinusitis |
| Severe infection  | Encephalitis, meningitis, myocarditis and cardiomyopathy, intestinal intussusception, disseminated infection, sudden infant death syndrome |
| Unknown           | Missing clinical information |
| Urinary tract infection | Dysuria, haematuria, haemorrhagic cystitis, and renal allograft dysfunction |
Results

Descriptive Epidemiology of HAdV Cases in Bern

We analysed samples that were referred to the IFIK for HAdV screening from January 1998 to December 2017. Of the 43,562 samples screened for HAdV from suspected patients, 34,556 (79.3%) samples were identified as unique patient cases based on patient I.D. and year of entry. Based on the laboratory confirmed results, 1302 (3.8%) were identified as positive. Of the 1302 HAdV positive cases, one case had information missing on age, gender, and clinical information, and was thus excluded from further analysis. The median age of HAdV positive patients was 3 years (range 0 months – 100 years; Table 2). A significant difference in the number of HAdV positive cases among males and females was observed ($P<0.001$) with females accounting for 42.4% (552/1301) of the cases. A significant difference in the frequency of positive HAdV cases among the different age groups was also observed ($P<0.001$), with the highest number of HAdV positive cases being found among young children aged ≤4 years (744; 57.2%), followed by those in the age group 20–44 years old (163; 12.5%). The least age group (59; 4.5%) was for children between 5 and 9 years. Logistic regression analyses confirmed that the overall effect of age category was statistically significant in determining positivity for HAdV infection as compared to those aged ≤4 years ($P<0.001$).

The yearly positive rate for HAdV ranged from 2.0% to 6.1%, with an average rate of 3.9%. A peak in proportion of HAdV positive cases was observed among young children aged ≤4 years (744; 57.2%), followed by those in the age group 20–44 years old (163; 12.5%). The least age group (59; 4.5%) was for children between 5 and 9 years. Logistic regression analyses confirmed that the overall effect of age category was statistically significant in determining positivity for HAdV infection as compared to those aged ≤4 years ($P<0.001$).

Table 2 Characteristics of the Study Population

| Demographic Characteristic | N (%) |
|----------------------------|-------|
| HAdV Positive Cases        | 1301* (100) |
| Gender                     |       |
| Female                     | 552 (42.4) |
| Male                       | 747 (57.4) |
| Unknown                    | 2 (0.15)  |
| Age Category (Years)       |       |
| 0–4                        | 744 (57.2) |
| 5–9                        | 131 (10.1) |
| 10–19                      | 59 (4.5)  |
| 20–44                      | 163 (12.5) |

Table 2 (Continued).

| Demographic Characteristic                  | N (%) |
|---------------------------------------------|-------|
| Sample Types                                |       |
| Upper respiratory tract                     | 731 (56.2) |
| Conjunctivitis swab                         | 243 (18.7) |
| Stool                                       | 184 (14.1) |
| Swab                                        | 59 (4.5)  |
| Lower respiratory tract                     | 48 (3.7)  |
| Other                                       | 19 (1.5)   |
| Genital swab                                | 7 (0.5)   |
| Blood                                       | 5 (0.4)   |
| Unknown                                     | 5 (0.4)   |

| Clinical Presentation and Comorbidities     |       |
| Respiratory tract infection                 | 388 (29.8) |
| Unknown/missing                             | 268 (20.6) |
| Ophthalmic infection                        | 212 (16.3) |
| Fever                                       | 165 (12.7) |
| Gastrointestinal infection                  | 93 (7.1)   |
| Immunosuppressed                            | 39 (3.0)   |
| Respiratory tract infection + Gastrointestinal infection | 30 (2.3) |
| Severe infection                            | 16 (1.2)   |
| Other infection                             | 16 (1.2)   |
| Respiratory tract infection + Immunosuppressed | 12 (0.9) |
| Respiratory tract infection + Ophthalmic infection | 11 (0.8) |
| Co-infection                                | 7 (0.5)    |
| Respiratory tract infection + Other infection | 6 (0.5)  |
| Respiratory tract infection + Co-infection  | 5 (0.4)    |
| Urinary tract infection                     | 4 (0.3)    |
| Gastrointestinal infection + Other infection | 4 (0.3)  |
| Respiratory tract infection + Urinary tract infection | 4 (0.3) |
| Gastrointestinal infection + Urinary tract infection | 3 (0.2) |
| Fever + Other infection                     | 3 (0.2)    |
| Oncological disease                         | 2 (0.2)    |
| Gastrointestinal infection +Immunosuppressed | 2 (0.2)  |
| Gastrointestinal infection + Severe infection | 2 (0.2) |
| Respiratory tract infection + Severe infection | 2 (0.2) |
| Ophthalmic infection + Co-infection         | 1 (0.1)    |
| Severe infection + Immunosuppressed          | 1 (0.1)    |
| Respiratory tract infection + Gastrointestinal infection + Other infection | 1 (0.1) |
| Respiratory tract infection + Gastrointestinal infection + Urinary tract infection | 1 (0.1) |
| Respiratory tract infection + Ophthalmic infection + Co-infection | 1 (0.1) |
| Respiratory tract infection + Ophthalmic infection + Gastrointestinal infection | 1 (0.1) |
| Respiratory tract infection + Ophthalmic infection + Other infection | 1 (0.1) |

Note: *The total of 1301 corresponds to 3.76% of the total suspected HAdV cases over the 20-year period.
occurred in years 2009 and 2010, with 110 and 112 cases, respectively (Figure 1A). A heat map of the monthly incidence showed that HAdV appeared to occur throughout the year. Although the data showed no clear seasonality, a pattern reminiscent of HAdV outbreaks was observed particularly in January 2010 (n = 24), July 2009 (n = 33), and March 1998 (n = 24). The mean expected incidence of HAdV cases per month from 1998 to 2017 was 5.43 cases (95% CI 1.86–12.26 cases). Therefore, HAdV incidence of ≥13 cases was defined as significantly higher number than the mean expected incidence (Figure 1B).

Upper respiratory tract samples were the most common positive sample types (N = 731; 56.2%), followed by conjunctivitis swab (243; 18.7%), and stool (184; 14.1%). HAdV was least detected from EDTA blood and genital swab samples. Higher numbers of positive conjunctivitis swab samples were observed in years 2009 and 2010 than in any other year (Figure 2A). Among young children, upper respiratory tract and stool samples were associated with the largest number of positive cases, whilst conjunctivitis swab samples were the most positive among young adults, middle aged and elderly (Figure 2B). A peak in HAdV cases was also observed between the years 2009 and 2010, as compared to other years in the study period, across all age categories. The most unexpected finding was a peak in the number of HAdV cases reoccurring every 4 years among children ≤4 years old, with increased incidences during the years 1998, 2001, 2002, 2006, 2010,

Figure 1 (A) Annual distribution of HAdV suspected cases and the overall percentage of HAdV positive cases per year referred to the IFIK from 1998 to 2017. (B) Monthly incidence of HAdV positive cases over the study period. The mean expected incidence of HAdV cases per month from 1998 to 2017 was 5.43 cases (95% CI 1.9–12.3 cases). Incidences larger than the upper CI (12) are indicated in bold font.
and 2014, with an average rate of 3.91% (Figure 3A and 3B).

**Molecular Typing of Cases Associated with Increased Incidence in Years 2009 and 2010**

Of the 222 HAdV positive samples, 154 (69.4%) samples were available for HAdV molecular typing. Out of the 154 samples subjected to typing, HAdV genotypes were identified in 145 samples, including 63 (43.4%) samples from patients aged 0–4 years, 8 (5.5%) from 5 to 9-year-olds, 5 (3.4%) from 10 to 19-year-olds, 21 (14.5%) from 20 to 44-year-olds, 21 (14.5%) from 45 to 64-year-olds, and 27 (18.6%) from 65 to 100-year-olds. Fourteen HAdV genotypes were identified to be circulating during this time period (Figure 4A), with 10 and 11 different HAdV genotypes identified in the years 2009 and 2010, respectively. The predominant circulating types in these 2 years were HAdV8 (63; 43.4%), HAdV2 (31; 21.4%), HAdV1 (21; 14.5%), and HAdV3 (11; 7.6%).
with HAdV species D genotypes accounting for the majority (48.3%) of the cases. However, in the year 2010 only, HAdV2 belonging to HAdV species C became the predominant genotype accounting for 29.9% (20/67) of the cases, followed by HAdV8 with 26.9%, and HAdV1 with 16.4% of the cases. Genotypes HAdV41, HAdV56, and HAdV64 were detected in the year 2009 only, whilst HAdV6, HAdV7, HAdV28, and HAdV53 were detected in the year 2010 only. Among samples from those aged 0–4 years (n = 63), HAdV2 (46.0%), HAdV1 (30.2%), HAdV3 (7.9%), and HAdV5 (6.3%) were the main genotypes. Among samples from 5 to 9-year-olds (n = 8), HAdV3 (50.0%) was the main genotype. HAdV8 was the predominant genotype, accounting for 40.0%, 71.4%, 85.7%, and 100% of the cases, among samples from 10–19, 20–44, 45–64, and 65–100-year-olds, respectively (Figure 4B).

HAdV cases with ophthalmic infection (n = 36), respiratory tract infection (n = 36), and with unknown/missing clinical information (n = 51), accounted for the majority of the genotyped samples (Figure 4C). Most cases presenting with ophthalmic infection (88.8%) and those with unknown/missing clinical information (74.5%) were identified to be infected with HAdV species D genotypes, particularly HAdV8. HAdV cases presenting with respiratory tract infection were associated with HAdV species C genotypes (HAdV1, HAdV2, and HAdV5), and HAdV species B genotype (HAdV3). Similarly, the former HAdV species C genotypes were also found in patients presenting with fever. Among patients presenting with gastrointestinal infection, HAdV2, HAdV3 and HAdV41 were identified. HAdV6 was identified in a case presenting with immune suppression. Among those who had more than one clinical presentation and comorbidity (ie, respiratory tract infection and gastrointestinal infection, respiratory tract infection and immune suppression, respiratory tract infection and co-infection), HAdV2 and HAdV3 were identified.

Figure 3 (A) Epidemiological patterns of HAdV cases overtime among different age categories. (B) Comparison of temporal patterns between absolute and relative numbers of HAdV cases. In panel B, the dotted line represents the overall average number of cases.
Molecular Epidemiology of HAdV Between 1998 and 2017 in Young Children

Out of the 745 identified positive samples from young children (≤4 years old) during years 1998–2017, 198 (26.6%) were available for typing. We identified a wide variety of HAdV genotypes circulating among young children during this period (Figure 5A), with HAdV1 (n = 49), HAdV2 (n = 93), HAdV3 (n = 17), and HAdV41 (n = 13) being the most common HAdV genotypes accounting for 24.7%, 47.0%, 8.6%, and 6.6% of the cases, respectively. Most typed samples were from the years 2010 (n = 38), 2009 (n = 25), 2006 (n = 24), 2002 (n = 15), 2001 (n = 17), and 2000 (n = 15).

We did not identify instances of co-infection with different genotypes of HAdV. The majority of HAdV genotypes were identified among those presenting with respiratory tract infections, fever, gastrointestinal infection, and missing clinical information (Figure 5B). HAdV2, HAdV5 and HAdV1 were identified in three dead young children (≤4 years old) in years 2006, 2012, and 2014, respectively. All three cases were associated with sudden infant death syndrome.

Phylogenetic Analysis

The nucleotide sequences of the partial hypervariable hexon gene 1–7 from HAdV genotypes identified in this study were
aligned and analysed with sequences from circulating HAdV genotypes identified in China including Taiwan, Egypt, France, Germany, Japan, Korea, Russia, Sweden, and the USA (Figure 6). The phylogenetic analysis showed that the sequences of the HAdV genotypes (HAdV1, HAdV2, HAdV3, HAdV4, HAdV5, HAdV6, HAdV7, HAdV8, HAdV28, HAdV37, HAdV41, HAdV53, HAdV56, and HAdV64) identified in the years 2009 and 2010 in this study from patients of all ages clustered with other international sequences of similar HAdV genotypes and exhibited similarity >98% at the nucleotide level. Similarly, phylogenetic analysis of the 14 HAdV genotypes identified among young children ≤4 years infected with HAdV from years 1998 to 2017 in our study clustered with international sequences of the same HAdV genotypes (Figures S1A and B).

Discussion

Our 20-year retrospective observational study from patients across all age categories underlines the need to carry epidemiological studies and molecular typing of HAdV among patients with HAdV infection. Although long-term epidemiological trends demonstrated no clear seasonality, a pattern reminiscent of recurring outbreaks was evidenced with high number of HAdV cases observed...
in January 2010, July 2009 and March 1998 as compared to other months or years during the study period. Genotyping of samples from years 2010 and 2009 identified HAdV8 as predominant genotype causing HAdV infection among young adults, middle-aged and elderly, and HAdV1 and HAdV2 as predominant genotypes causing HAdV infection among young children. Although HAdV infections are known to fluctuate over time, increased cases of HAdV infections were evidenced among young children every 4 years. This study is the first to report such recurring pattern. Previously, Heo and colleagues showed that cases of HAdV respiratory...
infection are common during winter and spring. A study by Yao and colleagues concluded that HAdV prevalence may be related to age and season.\textsuperscript{24} Our findings suggest that increased HAdV infection in Switzerland was related to age, but to a lesser extent to a particular season, as HAdV infections occurred throughout the year. Also when detected, HAdV was predominantly found among those aged \( \leq 4 \) years. HAdV is therefore confirmed to be an important pathogen in clinical practice, mainly in the paediatric population.

The high incidence of HAdV8 associated with ophthalmic infections among young adults, middle-aged and elderly in the years 2009 and 2010 is consistent with what was seen in Germany for instance.\textsuperscript{25} Our finding is also consistent with other studies that showed HAdV8 to be a major culprit of adenovirus-associated conjunctivitis.\textsuperscript{9,26,27} In addition, the predominance of HAdV species C genotypes, particularly HAdV1 and HAdV2, found among young children is also supported by other studies.\textsuperscript{28} As young children have developing immune system, they may be more prone to infections. In the same line of evidence, most HAdV infections in immunocompromised hosts are known to be associated with HAdV species C genotypes 1 and 2.\textsuperscript{14,29} Our study focused on molecular typing of HAdV positive samples collected between 1998 and 2017 among young children \( \leq 4 \) years in order to identify the genotypes associated with an increase of cases of HAdV infection among this age category every 4 years. Various genotypes were identified to be causing HAdV infections, among which HAdV1 and HAdV2 genotypes were predominantly detected in the years with reoccurring peaks. These genotypes were also observed in the non-reoccurring peaks, but to a lesser extent. These typing results give insight into circulating HAdV genotypes causing disease mainly among young children who are known to be at high risk of HAdV infection.

Although patients from all age categories may be susceptible to HAdV infections, more cases of HAdV infections in this study were observed among children \( \leq 4 \) years old. HAdV species C genotypes (HAdV1, HAdV2, and HAdV5) and HAdV species B genotype (HAdV3) were the most commonly identified genotypes, indicating that young children may be particularly sensitive to these genotypes. Previous studies have shown that HAdV infections in young children may be severe, including manifestations ranging from meningitis, intussusception, subacute focal encephalitis and myocarditis, to disseminated disease, or even death.\textsuperscript{30–33} Of particular note to this study are the five patients with HAdV infection who died in the 20-year period. Four patients who died were aged \( \leq 4 \) years (three males and one female) and one patient was a 6-year-old male. Our data showed that, of the three patients who died and whose samples were available for typing, one had been infected with HAdV1, one had been infected with HAdV2, and one had been infected with HAdV5. All three patients presented with sudden infant death syndrome. The case of another patient aged \( \leq 4 \) years who died suffering from disseminated adenoviral infection with hepatitis, but whose sample was not available in this study for typing, was reported by Steiner and colleagues.\textsuperscript{34} Although HAdV infections are usually self-limited and mild, several studies implicating HAdV causing fatal cases have been reported.\textsuperscript{5,6,35}

The association between HAdV genotypes and clinical presentations and comorbidities in infected patients have been examined in several studies: Wurzel and colleagues found that HAdV species C genotypes (HAdV1 and HAdV2) are predominant in the airways of young children and are therefore a likely etiological agent of the disease.\textsuperscript{36} Wang and colleagues showed that only HAdV B, C and E were identified among children with significant respiratory symptoms.\textsuperscript{37} Furthermore, Liu and colleagues reported that HAdV species B genotypes (HAdV3 and HAdV7) are predominant genotypes among paediatric population presenting with acute lower respiratory tract infections.\textsuperscript{38} A strong association of HAdV40 and HAdV41 with gastroenteritis is known. In our study, we noticed that, besides these genotypes, other HAdV genotypes were identified in those presenting with gastrointestinal infection (Figure 5B), a finding that is similar to what was reported by Kumthip and colleagues.\textsuperscript{39} In addition, it is known that specific HAdV species have distinct tissue tropism which may determine the clinical presentation.\textsuperscript{40}

The presence of HAdV in respiratory tract and stool samples from patients not presenting with any clinical symptoms might be detected incidentally, thus making interpretation of the clinical significance of a positive result difficult. Yet, in our study, samples of the patients who had any clinical presentation associated with HAdV infection were referred for laboratory diagnostics testing to our institute upon the request of the seeing physician when an adenovirus infection was suspected. Therefore, majority of the HAdV positive, hospitalized patients in this study had an adenovirus associated clinical presentation. Persistence of HAdV in a latent state among young children has been described.\textsuperscript{41–43} Reports of asymptomatic individuals shedding virus into the stool was also described.\textsuperscript{41,44,45} In our study, 86.1% of young children
≤4 years old whose stool sample tested positive for HAdV had a clinical presentation, with gastrointestinal infection, fever, and respiratory tract infection being the most predominant clinical presentations.

The phylogenetic analysis showed that the sequences of HAdV genotypes identified in our study displayed great similarity with nucleotide sequences from HAdV genotypes isolated in other countries. Although this finding may indicate that the same circulating genotypes may be found at the global scale, we cannot exclude that the partial sequences of the hypervariable hexon gene does not allow enough resolution among genotypes. As such, targeting other regions of the HAdV genomes such as fiber, penton base, and/or DNA polymerase genes may provide further information on differences among circulating HAdV genotypes, may help clarify the HAdV genotypes and also identify recombination events.

Our study has a number of limitations. First, our reported trends likely underestimate the true burden of HAdV infections in both the hospitalized patients and in the community, as samples sent for laboratory diagnostic testing to the IFIK rely on request by physicians, when an adenovirus infection is suspected. Second, samples were tested using different diagnostic methods for HAdV and some methods are known to be less sensitive for HAdV detection than others. Therefore, there is a chance that this could have also affected the results of our study. Third, some of the clinical information was missing, hence restricting interpretation of trends in HAdV clinical presentation associated with circulating genotypes. In addition, clinical information was obtained from electronic clinical records provided to the IFIK diagnostic department and could not be improved in most of the cases. Fourth, several HAdV positive samples could not be genotyped due to unavailability/lack of sample material. Although the missing information was not systematic for a particular age group, sample type or year, the unavailable information might have biased our results and therefore, the reported HAdV genotypes might not be fully representative of circulating HAdV genotypes in a given year. In addition, a complete quantitative description of the main genotype(s) causing an increase in HAdV infection every 4 years among young children could not be reached due to this lack of samples.

In conclusion, our study gives insight on HAdV infections from the largest hospital in Bern and other regional clinics serving the Swiss population. Patients of various age categories were identified to be infected, with peak years in 2009 and 2010. In these 2 years, HAdV8 was identified as the predominant genotype and was associated with HAdV infections among young adults, middle aged and the elderly, whilst HAdV2 and HAdV1 genotypes were identified to be predominantly causing HAdV infection among young children. The findings support that variation in the clinical presentation and epidemiology of HAdV genotypes exists, and may present unexpected patterns when long-term records are examined. By uncovering HAdV molecular epidemiology and its clinical presentations in our 20-year observation study, we have highlighted the existence of differences in HAdV infections among various age categories and circulating HAdV genotypes. Future work will be performed to sequence the whole genomes of representative HAdV isolates to assess the recombination potential in our strain collection.

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Disclosure
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