Genotypic diversity, circulation patterns and co-detections among rhinoviruses in Queensland, 2001

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Abstract

**Purpose.** Rhinoviruses (RVs) occur more frequently than other viruses and more often in people displaying symptoms than in those without. We sought to estimate the spectrum of RV diversity, RV species seasonality and to analyse RV involvement in respiratory virus co-detections.

**Methodology.** A convenience collection of 1179 airway sample extracts from patients with suspected respiratory infections, collected during 2001, was subjected to comprehensive molecular testing.

**Results.** RVs were the most common virus detected. We were able to genotype ~90% of RV detections, identifying 70 distinct RVs, spanning all three species. RV-Bs were under-represented. We found RV species co-circulated at times, although one species usually dominated. Each species displayed a bimodal distribution.

**Conclusion.** Notably, RVs and influenza A viruses (IFAV) seldom co-occurred, supporting their roles as primary pathogens of the airway among acutely ill infants. Whether RV circulation has a moderating or controlling effect on the IFAV season or is controlled by it cannot be determined from these data. Despite the frequent perception that RVs commonly co-occur with another virus, our findings indicated this was not always the case. Nearly 80% of RV detections occurred alone. Understanding more about population-level interference between viruses may allow us to harness aspects of it to generate a non-specific antiviral intervention that mimics a putative protective effect. For routine respiratory virus screening to best serve the patient, RV testing should be a principal component of any acute respiratory illness testing algorithm throughout the year.

INTRODUCTION

Rhinoviruses (RVs) are the largest related assemblage of genetically and antigenically distinct respiratory pathogens known, comprising 168 genotypes. These picornavirus species often occur in symptomatic young children from community and hospital populations where they create a sizable burden for management and are a frequent trigger of wheeze [1–3]. Picornavirus infections are more often observed among children presenting to an emergency department than other respiratory virus infections [4]. Nearly three-quarters of respiratory virus detection episodes during the first 28 days of life are due to RVs and over half are symptomatic [3]. Until recently, much of what is known of RV diversity, epidemiology and clinical impact was determined using human studies and cell culture methods in the 1950s to 1980s. In 2006, the molecular discovery of a genetic clade of RVs that could only be cultured in vitro using sophisticated air–liquid interface cultures was subsequently ratified as a third species, Rhinovirus C (RV-C) [5, 6]. To date, RV-C has added 56 distinct RV types to the genus Enterovirus, family Picornaviridae. As more is learned, conclusions reached by some earlier studies have required re-examination and confirmation [5–7].

We aimed to estimate the spectrum of RV genotypes, species seasonality and RV involvement in co-detections in...
Queensland using a convenience collection of airway sample extracts from patients with suspected respiratory infections, collected during 2001 and tested using molecular tools expected to account for all RV species.

**METHODS**

**Specimen extracts**

Specimen extracts (n=1179; 94% nasopharyngeal aspirates) for retrospective RV testing originated from an unselected sample of mostly young patients (57.5% male). Ages ranged from 1 day to 90.1 years (mean 5.4 years, median 1.5 years) and patients were located across Queensland, presenting to a hospital or clinic serviced by Pathology Queensland, Queensland Health (PQ). Patients had symptoms of acute respiratory infection during 2001. There was no systematic sampling protocol used, thus our sample did not represent the total population of extracts received by PQ for testing in 2001. Extracts included nasopharyngeal aspirates (85.5%), lavage (9.5%), endotracheal aspirates (2.2%) and swabs (1.3%). Nucleic acids had been previously extracted and stored at −80°C as previously described [8].

**Virus screening, RV species and type designation**

Extracts had been previously tested by PQ using direct or culture-amplified direct fluorescent assay to detect respiratory syncytial virus (RSV), adenoviruses (AdV), parainfluenza viruses (PIVs) and influenza viruses A and B (IFAV, IFBV) [9]. PCR was not in routine use at PQ in 2001. This study applied additional previously described real-time RT-PCR (RT-rtPCR) assays to detect human metapneumovirus (MPV) [8, 10] and human coronaviruses (HCoV) 229E, HKU1, NL63 and OC43 [11]. RVs were detected using a modified version of a previously described RT-rtPCR [12, 13]. The RV RT-rtPCR can also detect some human enterovirus (EV) types.

RV/EV-positive extracts were genotyped using a conventional nested VP4/VP2 RT-PCR assay incorporating previously described primers that include part of the 5′ untranslated UTR, viral protein coding regions VP4 and partial VP2 [14] no. 275; [15] no. 3252; [16] no. 528. First-round RT-PCR used OneStep RT-PCR (QIAGEN, Australia) or SensiFAST Probe No-ROX One-Step (Bioline, Australia) kits and subsequent second-round PCR used Bioline MyTaq HS DNA polymerase kit. In cases of failure to amplify a VP4/VP2 amplicon (~540 bp), part of the 5′ UTR (~400 nt) was amplified [16]. Amplicons were sequenced (BigDye sequencing kit v3.1, Applied Biosystems) and, after removal of the primer sequence (Geneious Pro v6) [17], RV and EV sequences were submitted to GenBank (accession numbers KF499366-KF499501, KF688607-KF688723). Curated and detailed methods are available online [18–20]. A virus variant was described as an ‘untypeable RV’ when a clean sequence could not be obtained from either genotyping assay, despite the extracts being repeatedly positive by RT-rtPCR assay. RV genotype was determined by best match using the following algorithm: when a query sequence returned BLAST comparison with ≥97% sequence identity in the 5′ UTR or ≥90% (for RV-B and RV-C) or ≥91% (for RV-A) in the VP4/VP2 region, with members assigned to a given type, our sequence was assigned as being a variant of that RV genotype.

**Statistical methods**

We provide descriptive statistics for these data, using counts and proportions. Following methods we described previously [21], univariate analysis was used to screen the relationships between picornavirus groups (RV-A, RV-B, RV-C or EV), season and demographic variables such as age and sex. For each virus, the probability of co-detection with another virus was assessed with Fisher’s exact test using 2×2 contingency tables. We used a threshold P-value of <0.05 for a statistically significant association. Logistic regression was used to estimate any effect of age, sex, season and other circulating virus types on detection of RV.

**RESULTS**

**Virus detections**

In 615 extracts (52.2% of those tested) at least one virus was detected. The most frequently detected virus or virus group was a picornavirus (Fig. 1 and Table 1; suspected RVs or EVs; n=296; 25.1% of all extracts, 48.1% of all virus positives) followed by RSV (n=101; 8.6% of all extracts), MPV (n=85; 7.2%). IFAV (n=76; 6.2%), PIV (n=66; 5.6%; 54 were PIV-3), AdV (n=37; 3.1%), HCoV-OC43 (n=26; 2.2%), HCoV-HKU1 (n=5; 0.4%), HCoV-NL63 (n=4; 0.3%) or HCoV-229E (n=1; 0.1%). No IFBV was detected and we did not seek human bocaviruses, influenza C virus or PIV-4.

**Demographic features of virus positives**

The virus-positive population had a median age of 1.5 years (IQR 0.06–3.5 years); this was lowest for RV and RSV cases (2.9 and 2.4 years, respectively). The age range varied among viruses in our population (Fig. S1, available in the online version of this article). Fewer than 5% of neonates or adults were RV positive (Fig. S2).

There are more females in the Australian population (0.9:1.0 [22]), but more males in the total patient sample (678/1179, 57.5%, P<0.0001 for an expected proportion of 50%). In the context of total patient extracts, 180/678 (26.6%) extracts tested from males were picornavirus positive compared with 116/501 (23.0%) of extracts from females (P=0.20). Similar results were seen in the context of the 615 virus-positive extracts. Of the 296 picornavirus positive extracts, 60.8% were from males and 39.2% from females, but this difference in proportion was comparable to the sex proportions in the picornavirus negative group and was not significant (P=0.32).

**RV speciation and genotyping**

A RV or EV type was assigned to 278 (93.9%) of 296 extracts (25.1%) that were picornavirus-positive by our screening RT-rtPCR. Most picornaviruses were RV-As and RV-Cs.
Table 1). In total, 70 distinct RV types circulated at our site, including variants of four types that could not be assigned to an existing genotype (Table S1). Twelve genotypes represented 50.7% of those found and three genotypes belonged to the ‘minor group’ of receptor-classified RVs. There are 32 known RV-B types therefore 19.2% of RV detections were expected if RVs are distributed randomly. However, only 3.7% of picornaviruses were RV-Bs, which is a significant under-representation (P<0.001). To ensure both genotyping methods performed as expected, we determined both 5′ UTR and VP4/VP2 sequences for ten RV positives and typing assignments agreed between targets using our thresholds.

**Seasonality**

Total virus detections did not present with any clear seasonality but individual viruses and virus groups did. Most detections were in spring and autumn (61.2% of positives; Figs 1 and 2). RSV detections peaked in autumn, PIVs, HMPV and HCoV-OC43 in spring, AdV and IFAV during late winter and EVs during summer and autumn.

Inspection of epidemic curves suggested some relationships between viral seasons. Spearman correlations supported these impressions, showing that both RV and RSV seasons were negatively associated with the IFAV season (Table S2). Positive correlations included RV with RSV; EV with RSV; AdV with PIVs; HMPV with IFAV, HCoV-OC43, HCoV-NL63 and PIV.

RVs were detected in all seasons, but a bimodal distribution was evident with peaks in spring (36.5% of all picornaviruses) and autumn (35.8%) and a trough in winter (13.2%; summer extract numbers were low, so prevalence was difficult to determine; Fig. 2). We observed seasonal exchange between RV species; RV-As and RV-Cs swapped dominance throughout the year. Although their numbers were lower, RV-Bs (none detected in 4/12 months) and EVs (none detected in 7/12 months) exchanged peak positions of prevalence throughout the year.

**Analysis of co-detections**

Most viruses occurred as single detections (n=539; 87.6% of virus positives; Fig. 3). Co-detections were found in 76 extracts; 71 with two viruses detected and five with three viruses detected. No extract had more than three viruses identified. Viral co-detection totals also exhibited a bimodal peak (Fig. S3).

Associations of RVs with other viruses, sex and season, were examined using logistic regression models, and results were consistent with those from the univariate analyses (Table S3).

Detection of more than one virus occurred in 76 extracts. RVs were the most frequently detected viruses overall and were involved in the greatest total number of co-detections (n=60; 20.3% of all 296 RV-positive extracts) (Fig. 3; Table S4). Most (n=44; 73.3%) RV co-detections occurred in those 2 years of age or younger, and none of the 60 extracts originated from adults (>14 years of age). The proportion of each RV species
involved in co-detections differed ($P<0.0001$). Of the RV-As, 21 of 142 (14.8%) were co-detections while 7 of 11 (63.6%) RV-Bs and 22 of 112 (19.6%) RV-Cs were co-detections. One of the 12 EVs (8.3%) was a co-detection. RV-Bs were more likely to occur as co-detections than RV-As ($P=0.001$), RV-Cs ($P=0.003$) or EVs ($P=0.009$). Ten of the 19 (52.6%) untypeable picornaviruses were co-detections. Most RV co-detections were with paramyxoviruses (MPV, $n=21$; RSV, $n=17$; PIV, $n=16$) followed by AdV ($n=5$), IFAV ($n=2$) and HCoV-HKU1 ($n=1$). There were no co-detections identified between RV or EV species ($P<0.004$ for all comparisons) except for RV-B and EV. EV co-detections were rare overall; given a positive detection of EV, 0/11 (0%) extracts were positive for RV-B compared with 12/285 (4.2%) extracts negative for RV-B, $P=1.0$.

It was important to look at co-detections as a proportion of each virus, not just RVs. Other viruses were involved in a similar or greater proportion of co-detections compared to the RVs (Fig. 3). IFAV had one of the lowest proportions of

### Table 1. Features of picornavirus positive extracts.

|                      | RV-A     | RV-B     | RV-C     | EVs     | Unknown PV type | $P$-value for difference within PVs |
|----------------------|----------|----------|----------|---------|-----------------|------------------------------------|
| No. of PV detections total | 142 (91/51, 35.9) | 11 (3/8, 72.7) | 112 (71/41, 36.6) | 12 (5/7, 58.3) | 19 (9/9, 50.0) | 0.09                               |
| $n=296$ (no. of M/F, % female) | 296 (190/106, 64.7) | 11 (3/8, 72.7) | 112 (71/41, 36.6) | 12 (5/7, 58.3) | 19 (9/9, 50.0) |                                      |
| Expected % of RV species | 47.6 (80/168) | 19.0 (32/168) | 33.3 (56/168) | NA | NA |                                      |
| % of RV types from that species | 38.8 (31/80) | 15.6 (5/32) | 53.6 (30/56) | NA | NA |                                      |
| % positive, of total extracts (1179) tested | 12.0 | 0.9 | 9.5 | 1.0 | 1.6 | <0.0001 |
| % of virus-positive extracts (615) | 23.2 | 1.8 | 18.3 | 2.0 | 3.1 |                                      |
| Median, IQR age in years | 1.09, 0.41–2.35 | 1.92, 0.77–5.37 | 1.65, 0.95–2.63 | 3.34, 1.54–9.14 | 1.62, 0.51–7.10 | 0.02 |
| Mean age (range), years | 2.7 (0.03–26.1) | 3.0 (0.6–6.7) | 2.9 (0.04–37.9) | 6.3 (0.2–16.4) |                                    |

EV – enterovirus; IFAV – influenza A virus; IQR – interquartile range; NA – not applicable; PV – picornavirus; RSV – respiratory syncytial virus; RV – rhinovirus

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Fig. 2. Number of extracts and virus detections. The total number of extracts tested (solid line, right $y$-axis) and the number of any virus detections (broken line, right $y$-axis) as well as the number of each RV species or EV detected (bar graph, left $y$-axis), per month, 2001. The predominating RV species is described at the top and key seasons along the bottom. DOI: 10.6084/m9.figshare.6388850.
co-detections (9.6%) as did the EVs (8.3%) while HCoV-NL63 was not involved in any co-detections. RVs were significantly less likely than expected to be co-detected with IFAV ($P<0.0001$; Table S4). PIVs were less likely to be detected with RSV ($P=0.05$) or IFAV ($P=0.03$), and RSV was less likely to be found with IFAV ($P=0.002$) than expected.

**DISCUSSION**

Seventy distinct RV genotypes circulated among this symptomatic convenience population. Levels of genotypic diversity vary among other studies; some are higher than we observed, such as during the first year of life among the childhood origins of asthma cohort [23], but many observe fewer distinct genotypes [24–26]. Studies which do not account for RV species or genotype risk overestimating the length of RV shedding, falsely assuming persistence rather than sequential infection [27, 28]. Among our sample, RVs dominated in prevalence compared to all other respiratory viruses across most of 2001, notably waning during the seasonal IFAV epidemic.

Most picornavirus-positive extracts were genotyped, a fact that agreed with some studies, but not others [23, 28–32]. Lower rates of genotyping success in community studies may be due to lower viral loads among less severely ill subjects. We propose that our combination of real-time RT-PCR screening and conventional nested-PCR genotyping methods are an effective starting point for genotyping ~90% of RVs among ill populations, even from extracts stored for over a decade. Nonetheless, the presence of co-detections specifically among the RVs and EVs may have been underestimated by use of broadly specific detection and sequencing assays.

Others have found RV-Cs to be the dominant RV species in hospital-associated asthma exacerbations [1, 33]. We have previously found them to be numerically dominant but RV-A to be more clinically significant in non-hospitalized asthmatics [34]. We observed RV-B detections to be a significantly smaller proportion of total RV detections than was expected. We have previously noted this in studies of hospitalized populations [29, 34]. It is not clear why RV-Bs are under-represented. Perhaps the RV-Bs are less severe pathogens [23, 26] or their infections contribute disease to a different population than that tested here. It is noteworthy that RV-Bs are also under-represented in community populations [28, 35].
While the RV species often co-circulated during 2001, one species usually predominated each month, changing from month to month. Each species displayed bimodal peaks in autumn and spring and a trough in winter, which aligned with IFAV circulation. Exchange of prevailing species and turnover of prevalent genotypes is supported by other studies that employ molecular typing and span 12 months or more [15, 25, 36–39]. Shorter cross-sectional studies may misrepresent association between disease severity and RV species or genotype by failing to capture year-to-year variation in RV diversity.

Whether RV circulation has a moderating or controlling effect on the IFAV season or is controlled by it, cannot be determined from these data however these viruses seldom seasonally co-occurred. We and others have seen this and related interactions, termed virus interference, in other populations [29, 40, 41]. Virus interference is a consistent feature that some have attributed to RV dominance [42, 43]. It remains unclear how this apparent interference pattern is affected by the mostly paediatric and symptomatic population, the high proportion (86%) of nasopharyngeal extracts and relatively few swabs used in this study, whether influenza type, subtype or genotype plays a role, if pre-existing RV immunity can influence interference or whether certain RV strains or species play a disproportionate role in interference. Influenza may control a population’s susceptibility to other respiratory viruses, blocking their spread because of a strong population-level innate immune response generated by influenza strains during an annual epidemic ‘flu season’. Study of additional years may address these questions.

Initiation of seasonal influenza epidemics is mostly associated with rate of influenza virus change, strain replacement, availability of susceptible hosts and climatic conditions [44, 45]. Further study of virus interference is warranted and could inform our understanding of human respiratory diseases that exhibit temporal trends such as exacerbations of asthma and chronic obstructive pulmonary disease, otitis media and allergic rhinitis. Understanding a mechanism for interference may allow us to harness it to generate a non-specific antiviral intervention that mimics this putative protection.

We found nearly five times more co-detections in our study than were reported from among the first viral detections of We found nearly five times more co-detections in our study than were reported from among the first viral detections [3, 27]. EVs were observed to have an especially low proportion of co-detections in our 2001 sample. As reported in our earlier studies of populations from different years and Australian locations, we again observed that detection of an IFAV, EV, RV or RSV occurred with reduced likelihood of co-detection of any other virus in that patient’s sample [21, 29]. There was a higher likelihood of co-detection among extracts positive for an AdV or MPV. Numbers were small for the other viruses. The accepted understanding of IFAV directly associated with severe illness was supported here by a low proportion of co-detections.

Infants from 1 to 12 months of age were most likely to be affected by RV or RSV in our convenience sample population; this agreed with findings from the Brisbane community cohort [3]. RV-positive children were second youngest after RSV-positive children while IFAV-positive children were generally older than those positive for any other virus. Interestingly, it is often children younger than 2 years who are affected by severe influenza [46]. Most RV detections (single or multiple) occurred in infants, the largest contributing population. It is unknown whether the viruses detected in this population reflect those that concurrently circulated among mild illness or subclinical community infections during 2001.

This analysis provides useful comparative baseline data for ongoing and future analyses [28, 47]. There is no denominator from which to determine the rates of virus infection nor is there a specific clinical definition of each infected person available. Neither clinical impact nor admission status was sought in this study, but sampling was due to a clinical need. The small number of extracts available from January and February (summer) limited the strength of the RV analyses during this period. Also, the absence of testing for bocaviruses may have reduced the positive associations noted among the respiratory viruses because bocaviruses are frequent co-detections in extracts positive for more than one respiratory virus [5, 21].

Despite the frequent perception that RVs commonly co-occur with another virus, our findings here and elsewhere indicate this is less likely to occur for RVs than for other respiratory viruses [21, 29]. That nearly 80% of RV detections occurred in the absence of another virus once again supports the assertion that RVs are primary pathogens of the airway among acutely ill infants. If a respiratory tract sample is to be collected for any testing and if routine screening of respiratory viruses is to serve its patients best, RV should be a principal component of any acute respiratory illness testing algorithm throughout the year, with allowance for their relative replacement by IFAV cases during winter.

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Author contributions
IMM – funding, conceptualisation, investigation, visualisation, original draft preparation, review and editing, supervision; KEA – investigation, manuscript review and editing; CTYW – investigation, manuscript review and editing; RMG – formal analysis, manuscript review and editing; IMM planned the study, conducted experiments, analysed data, acquired funding and wrote the manuscript; KEA conducted experiments, analysed data; RMG performed statistical analyses and co-wrote the manuscript. All authors reviewed, revised and approved the final version.

Conflicts of interest
The authors declare that there are no conflicts of interest.
Ethical statement
This study was approved by the Queensland Children’s Health Services human research ethics committee (#HREC/06/QCHRH/097) and UQ medical research ethics committee #200900039.

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