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Predicting the Next Eye Pathogen: Analysis of a Novel Adenovirus


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ABSTRACT For DNA viruses, genetic recombination, addition, and deletion represent important evolutionary mechanisms. Since these genetic alterations can lead to new, possibly severe pathogens, we applied a systems biology approach to study the pathogenicity of a novel human adenovirus with a naturally occurring deletion of the canonical penton base Arg-Gly-Asp (RGD) loop, thought to be critical to cellular entry by adenoviruses. Bioinformatic analysis revealed a newly highly recombinant species D human adenovirus (HAdV-D60). A synthesis of in silico and laboratory approaches revealed a potential ocular tropism for the new virus. In vivo, inflammation induced by the virus was dramatically greater than that by adenovirus type 37, a major eye pathogen, possibly due to a novel alternate ligand, Tyr-Gly-Asp (YGD), on the penton base protein. The combination of bioinformatics and laboratory simulation may have important applications in the prediction of tissue tropism for newly discovered and emerging viruses.

RESULTS The HAdV-D60 penton base gene has a unique deletion of the hypervariable RGD loop. Viral cultures were obtained by nasopharyngeal swab from a male child with bronchiolitis. Initial testing suggested a mixed infection with adenovirus and human metapneumovirus. Limited genomic sequencing of the adenovirus suggested a unique adenovirus type. Subsequent whole-genome sequencing, phylogenetic analysis, and viral neutralization assays for the adenovirus in the specimen revealed the novel adenovirus to be a recombinant of several human adenoviruses, with sequence contributions from HAdV-D20 (hexon gene) and HAdV-D37 (penton base gene) but with a unique fiber gene (Fig. 1A; see also Fig. S1 to S3 in the supplemental material). Thus, this new human adenovirus species D type (HAdV-D) was designated HAdV-D60, based on current typing criteria (19). Unexpectedly, sequence analysis revealed deletion of the Arg-Gly-Asp

Infection entails complex and functionally integrated relationships between pathogen and host (1, 2). The synthesis of genomic, bioinformatic, cellular, and molecular methodologies permits a systems biology approach to predict the emergence of new and severe infectious agents. Human adenoviruses (HAdV) are ubiquitous, cause infections in the respiratory, gastrointestinal, genitourinary, and ocular mucosas, can be opportunistic in the immune-compromised host, and have been associated with obesity (3–8). Adenoviruses also are common choices for gene therapy vectors (9, 10). Thus, while adenoviruses continue to cause significant morbidity and mortality in the human population, their existence also provides a potential benefit for the treatment of patients with an even broader range of ailments.

Since the first adenovirus was characterized in 1953 (11, 12), over 60 human adenoviruses have been recognized as unique types. Analysis of whole-genome sequence data for existing and new HAdVs confirmed a critical role for homologous recombination in adenovirus evolution, leading to new and sometimes serious human infections (13–18). The emergence of new HAdV types, with several associated with severe eye infection (13, 16, 18), prompted us to apply a systems biology approach to try to predict the ocular tropism of a previously uncharacterized and highly novel HAdV, isolated by nasopharyngeal swab from a 4-month-old boy with severe bronchiolitis. A combined genomic, bioinformatic, and biological analysis identified a unique deletion in a key protein of the viral capsid and further suggested the potential of this virus to cause significant ocular infection. These results point toward a possible approach for predicting pathogenicity for newly identified and recently emergent human pathogens.
loop (amino acids [aa] 294 to 319) in the predicted penton base protein from a representative type from each HAdV species. The highly conserved RGD sequence is bolded and boxed in red. (C) SimPlot analysis of the HAdV-D60 penton base gene queried against all completely sequenced HAdV-D penton base genes. The location, in relation to the nucleotide sequence, of the two major hypervariable loop (HVL1 and RGD) domains of the protein are presented above the graph. HAdV-D37 sequence similarity is presented in red. (D) Predicted protein model of the HAdV-D60 penton base protein based on homology modeling of the crystal structure of HAdV-C2. The absence of the RGD loop in HAdV-D60 is shown in red (arrow). The RGD loop for HAdV-C2 was not resolved in crystal structure and was modeled and added (arrow and in red).

Viral replication (Fig. 2B) and E1A transcription (Fig. 2C; Fig. S6) for HAdV-D60 also occurred but were delayed compared to those for HAdV-D37. Together, these data demonstrate that HAdV-D60 infects and replicates in ocular surface epithelial cells, albeit more slowly than HAdV-D37.

HAdV-D60 infects the cornea in vivo and is hyperinflammatory. Since in vitro infection of ocular surface epithelial cells was consistent with corneal tropism, we examined the capacity of HAdV-D60 to induce corneal inflammation in vivo by testing various titers of the virus in an adenovirus keratitis mouse model (22, 23). Surprisingly, HAdV-D60 induced threshold clinical inflammation at an infectious dose considerably (~100-fold) lower than that for HAdV-D37 (Fig. 2D; see also Fig. S7A in the supplemental material). By flow cytometry, roughly 4-fold more infiltrating CD45+ cells infiltrated the HAdV-D60-infected cornea than infiltrated corneas with similar titers of HAdV-D37 (Fig. 2E). Consistent with increased infiltration by leukocytes, cytokine levels of interleukin 6 (IL-6), CXCL1, CXCL10, alpha interferon (IFN-α), and IFN-β in the corneas of HAdV-D60-injected mice were significantly elevated compared to those in mice with control HAdV-
D37 infection (Fig. 2F; Fig. S7B). These results show that HAdV-D60 capably induces keratitis in the mouse.

**HAdV-D60 contains a novel potential integrin binding motif.** We previously showed that administration of an RGD 15-mer identical to the RGD loop in HAdV-D37, when coinjected with HAdV-D37, reduced resultant inflammation compared to a KGE-containing control 15-mer otherwise identical to the RGD peptide (23). Because HAdV-D60 lacks the penton base protein RGD motif thought essential for viral internalization via binding to and aggregation of host cellular integrins, we predicted that a similar experiment performed with HAdV-D60 would show no suppression of corneal inflammation by either the RGD 15-mer or its KGE control. Surprisingly, administration of the RGD 15-mer identical to the RGD loop in HAdV-D37, when coinjected with HAdV-D60, reduced resultant inflammation compared to the KGE containing control (Fig. 3A), suggesting that integrin binding remained an important element in the pathogenesis of infection or, less likely, suggesting a direct effect of the RGD ligand on the immune response (24). Protein modeling of the HAdV-D60 penton base illustrated the RGD loop deletion, but also identified a Tyr-Gly-Asp (YGD) motif (Fig. 3B), on the outer upper region of the modeled penton base protein. This motif was found to be conserved in all known HAdV-D penton base genes, suggesting novel and redundant strategies for penton-mediated integrin binding and aggregation (25). Notably, a YGD 15-mer identical to the amino acid sequence surrounding YGD in the HAdV-D60 penton base protein prevented binding of HAdV-D60 to cultured human corneal fibroblasts (Fig. 3C). Previous work demonstrated a key role for downstream expression of CXCL8 in human adenovirus keratitis (26). Treatment of human corneal fibroblasts with the YGD 15-mer also reduced CXCL8 expression in HAdV-D60 infection (Fig. 3D) and abrogated clinical and histological keratitis in the HAdV-D60-infected mouse cornea compared to those in the KGE control (Fig. 3E), suggesting a possible role for the YGD motif on the penton base. No other potential integrin ligand was found on the HAdV-D60 penton base protein.

**DISCUSSION**

Here, we report the identification of a novel human adenovirus and possible bronchial pathogen, HAdV-D60, which is partially recombinant with HAdV-D37, a known eye pathogen, and we describe an approach to test its potential to cause severe eye infections. Whole-genome sequencing revealed a recombinant genome with a recombinated and RGD loop-deleted HAdV-D37 pen-
This was surprising, as the RGD or similar motif is thought to enable viral entry via host cellular integrins for all human adenoviruses (20, 27). HAdV-D60 represents the first reported HAdV within species D lacking the specific RGD motif, and this is the first report of any HAdV missing the RGD loop in its entirety. Notably among previously known HAdVs, HAdV-F40 and HAdV-F41 express an external penton base loop with a modified integrin binding motif (28, 29); the loop on HAdV-F40 penton base protein expresses RGAD, while HAdV-F41’s penton base loop has YGD.

The capacity of HAdV-D60, with a penton base protein similar to that of HAdV-D37, to infect the ocular surface epithelium and induce keratitis despite the absence of an RGD loop suggests that other components of the penton base protein may play a role in tissue tropism. It is known that the fiber protein (see Fig. S3B in the supplemental material), in particular, the fiber knob, is an important determinant of tropism for the ocular surface epithelium. A single amino acid change in the knob domain of the fiber protein was implicated in conjunctival tropism (30). The fiber of HAdV-D60 was most similar to that of HAdV-D20, a virus not known to infect the ocular surface. Therefore, we predict effects on viral tropism also by other viral components beyond the fiber and penton base. Interestingly, in the absence of the RGD-containing penton base loop, HAdV-D60 infection of conjunctival and corneal cells in vitro appeared to be slowed relative to that with HAdV-D37 and viral gene expression was considerably delayed. Thus, the RGD loop appears to be important but dispensable for viral entry, possibly due to a surrogate YGD integrin binding motif on the virus. It is intriguing that HAdV-F40 and -F41, which are relatively fastidious enteric viruses (28, 31), also lack the penton base protein RGD motif and express the penton base YGD motif (28, 29). It is unknown at present whether the YGD on the adenovirus penton base protein is exposed sufficiently to interact with cellular integrins during virus attachment and entry. Crystalization of the HAdV-D60 penton base protein is in progress and may help resolve this question. Alternatively, free or altered penton base capsomer due to partial virus uncoating at the cell surface may aggregate cellular integrins and assist in viral entry after fiber knob binding (32, 33). Future work is needed to identify whether YGD represents a novel and redundant motif for adenovirus internalization. If inflammation is initiated by virus binding to its cellular receptor(s), then perhaps delayed internalization by...
HAdV-D60 is, in part, responsible for its substantially greater inflammatory capacity in the cornea in vivo.

Our data suggest that HAdV-D60 may cause ocular infection, but our synthesis can only predict, not foretell with certainty. Further, it is unclear to what degree HAdV-D60 is currently or might in the future become prevalent in the community. At present, the population distribution of HAdV-D60 is unknown, but another virus identical to HAdV-D60 with the exception of a recombined E3 transcription unit was recently isolated from the gastrointestinal tract of an HIV-infected patient in England (GenBank accession number JN162672) (34), suggesting that genome variants of HAdV-D60 may be in circulation outside Canada. Whole-genome sequencing recently identified 10 new or previously unidentified HAdV types, and surprisingly, three of these were isolated from patients with epidemic keratoconjunctivitis (13, 16, 18). Until or unless HAdV-D60 is identified in an outbreak of ocular infection, the true ocular pathogenicity of HAdV-D60 will remain unknown. The synthesis of in silico, in vitro, and in vivo experimental approaches, here applied to a novel virus in a hypothesized ocular infection, represents one potential means to predict tropism and pathogenicity for new or emerging infectious agents before they cause serious outbreaks of disease.

MATERIALS AND METHODS

Virus, cells, and animals. HAdV-D60 (GenBank accession number HQ007053) was cultured from a 4-month-old male infant with broncholitis in Edmonton, Canada. Respiratory virus testing was performed at the Provincial Laboratory for Public Health, Calgary, Canada. Nasopharyngeal swab test results for influenza A virus, influenza B virus, parainfluenza (types 1, 2, 3, and 4), and respiratory syncytial virus A and B were negative, but testing results for adeno virus and human metapneumovirus were positive. Blood and urine cultures were negative. HAdV-D60 was cultured directly from a nasopharyngeal swab using A549 cell culture and identified as a potentially novel virus on the basis of limited Sanger sequencing. Subsequent whole-genome sequencing of the virus capsid hexon gene. Subsequent whole-genome sequencing led to typing of the virus as HAdV-D60, the next available type number in GenBank, on the basis of its unique genome and phylogenetic analysis, as approved by the Human Adenovirus Working Group (http://hadvwg.gmu.edu/) (19), a collaboration between the adeno virology community and the National Center for Biotechnology Information (NCBI, Bethesda, MD). HAdV-D37 (GenBank accession number DQ090090) was obtained from the American Type Culture Collection (ATCC, Manassas, VA). For use in these studies, HAdV-D37 and HAdV-D60 stocks were grown in A549 cells and purified by a cesium chloride gradient.

Human lung carcinoma cell line A549 (CCL-185), a human alveolar epithelial cell line that was previously shown to support HAdV virion production (35), was obtained from ATCC. Telomerase-immortalized human corneal epithelial (TKE) cells were the kind gift of Jerry Shay (University of Texas—Southwestern Medical Center) and were cultured in defined keratinocyte serum-free medium with supplements (Gibco, Grand Island, NY). Human conjunctival epithelial (Chang) cells were obtained from the ATCC. Primary human corneal fibroblasts (HCF) were isolated from human donor corneas as previously described (26). Cells were maintained in Dulbecco’s modified Eagle’s medium containing 10% heat-inactivated fetal bovine serum. Eight- to 12-week-old wild-type female C57BL/6J mice were purchased from Jackson Laboratories (Bar Harbor, ME).

The research described in this study conformed to the tenets of the Declaration of Helsinki. The virus studied was cultured from a human patient as part of routine clinical care, deemed necessary by the clinical provider. The virus isolate would otherwise have been discarded and was deidentified for analysis and was therefore determined to be exempt from review by the Human Studies Committee at the Massachusetts Eye and Ear Infirmary, a Harvard Medical School affiliate. All animals used in this study were treated humanely and in full agreement with the Association for Research in Vision and Ophthalmology resolution on usage and treatment of animals in research. Approval for the use of animals in this study was obtained from the Massachusetts Eye and Ear Infirmary Animal Care Committee.

Genome sequencing, annotation, and analysis. Purified HAdV-D60 DNA was sequenced on a Roche 454 DNA sequencer (Branford, CT) by Eurofins MWG Operon (Huntsville, AL). Gaps, ambiguous sequences, and the inverted terminal repeats were sequenced using an ABI 3130xl (Applied Biosystems, Carlsbad, CA) via PCR amplification. Quality control included sequence annotation and comparison with HAdV genome landmarks. Data were assembled using CLC Genomics software (CLC bio; Cambridge, MA), with an N50 average of 5,260. Minimal sequence coverage was eight-fold (Sanger chemistry) and 17- to 20-fold (Next Gen), with the Next Gen data noted by Operon at an accuracy of greater than 99% (Q20 or better).

The software programs mVISTA Limited Area Global Alignment of Nucleotides (LAGAN; http://genome.lbl.gov/lavan/submit.shtml) and Molecular Evolutionary Genetics Analysis (MEGA) version 4.0.2 (http://www.megasoftware.net) were used for global pairwise sequence alignment and phylogenetic analysis as previously described (14, 15, 18, 36–41). For phylogenetic trees, branches with bootstrap values below 70 (indicative of low confidence) were collapsed. DNA sequence recombination analysis was performed with SimPlot 3.5.1 software (http://sravy.med .som.jhmi.edu/SCRoftware/simplot/), including the options of Bootscan and SimPlot analysis, to identify possible recombination sites (42). Default settings were used, and HAdV-D60 was used as the reference sequence for analysis.

Protein analysis and modeling. Multisequence amino acid alignment was performed using CLC sequence viewer 6 (http://www.clcbio.com /index.php?id=28). A model of the HAdV-D60 penton base was constructed using Swiss-Model (http://swissmodel.expasy.org/) for homology modeling (43, 44). The crystal structure of HAdV-C2 (45) (Protein Data Bank [PDB] no. 1X9T) was used as a template for the model. The identity between the HAdV-C2 penton base and the HAdV-D60 penton base was 77%. The model generated by Swiss-Model was evaluated by PROCHECK (http://www.ebi.ac.uk/thornton-srv/software/PROCHECK/) and a Ramachandran plot, which identified 82.7% of residues in most-favored regions and 14.9% in additional allowed regions. To model the missing RGD loop from the crystal structure of the HAdV-C2 penton base, the RGD loop consisting of 78 amino acids was modeled using the Robetta Server (http://robetta.bakerlab.org/). Robetta provides both ab initio and comparative models of protein domains and uses the Rosetta fragment insertion method. Five PDB files were generated for the structure of the hypervariable RGD loop of HAdV-C2. One model was chosen and fit by DeepView software (http://spdbv.vital-it.ch/) on the crystal structure of HAdV-C2 to get the full penton protein. UCSF Chimera software (http://www.cgl.ucsf.edu/chimera) (46) was used for visualization and analysis.

Experimental infections. Cells grown to 80 to 90% confluence were infected in triplicate with virus at a multiplicity of infection (MOI) of 5 or mock infected with virus-free dialysis buffer, and virus-infected cells and/or supernatants were harvested at indicated time points and assayed, as described below. For some infections, HAdV-37 and HAdV-D60 were conjugated with Cy3 dye (GE Healthcare, Piscataway, NJ) according to the methods of Leopold et al. (47). One milligram of Cy3 dye was reconstituted in 1 ml of 0.1 M sodium bicarbonate (pH 9.3). Labeling was performed by conjugating Cy3 dye to virus at a concentration approximately equal to 1012 ADV particles/ml, where reconstituted Cy3 dye was 20% of the final solution. The mixture was allowed to incubate for 30 min in the dark with gentle mixing every 10 min, followed by overnight dialysis to remove the excess Cy3 dye.

Mice were anesthetized by intramuscular injection of ketamine (85 mg/kg of body weight) and xylazine (14 mg/kg). Anesthetic drops
Inhibition of infection by adenoviral penton base peptides. Synthetic 15-mer peptides were obtained from GenScript Corporation (Piscataway, NJ). The sequence of the HAdV-D37 wild-type penton base peptide including the RGD loop was DAVPRGDNYASAAEA, and the negative-control peptide with KGE in place of RGD, was DAVPKGNYASAAEA. The sequence of the HAdV-D60 wild-type penton base peptide including YGD was LSYTKGEPEKGVQSW. A unifying view of 21st century systems biology. FEBS Lett. 2009:135–167.

ELISA. Human cells were infected in triplicate with either HAdV-D60, HAdV-D37, or virus-free dialysis buffer and supernatants harvested for an enzyme-linked immunosorbent assay (ELISA) at 4 h postinfection for protein detection. Mouse corneas were removed at 16 h postinfection (n = 3/time point/group) and flash frozen in liquid nitrogen. Corneas were then homogenized in 400 μl of PBS with 1 mM phenylmethylsulfonyl fluoride (PMSF), 1 μg/ml aprotinin, and 10 μg/ml leupeptin (Sigma-Aldrich, St. Louis, MO). The lysates were centrifuged at 10,000 × g for 10 min at 4°C, and the supernatants were used for analysis. ELISAs for human CXCL-8 and mouse CXCL1, IL-6, CXCL10, IFN-α, and IFN-β (all from R&D Systems, Minneapolis, MN) were performed according to the manufacturer’s instructions. Each sample and standard was analyzed in duplicate. The plates were read on a microplate reader (Molecular Devices, Sunnyvale, CA) and analyzed (SoftMax software; Molecular Devices).

Flow cytometry. Corneas were dissected from mouse eyes at 4 days postinfection and cut into small (1- to 2-mm-diameter) fragments for subsequent digestion with 1 mg/ml collagenase type 1 and 0.5 mg/ml DNase (Sigma Chemical Co., St. Louis, MO) (13, 23). Single-cell suspensions were washed twice (300 × g, 5 min/wash) in PBS and then incubated on ice for 15 min with 2 μl anti-mouse Fc block (BD Pharmingen, San Diego, CA) in a total volume of 100 μl PBS-1% BSA. Following incubation, cells were centrifuged (300 × g, 5 min) and resuspended in 5% normal rat serum (Jackson ImmunoResearch Inc., West Grove, PA) for an additional 15 min on ice. Cells were then labeled with 4 μl anti-mouse fluorescein isothiocyanate (FITC)-conjugated anti-CD45 (clone 30-F11; BD Pharmingen, San Diego, CA) and incubated in the dark on ice for 30 min. Following incubation, the cells were washed 3 times with PBS-1% BSA (300 × g, 5 min/wash) and resuspended in PBS containing 1% paraformaldehyde. After overnight fixation at 4°C in the dark, cells were pelleted and resuspended in PBS-1% BSA and flow cytometry was performed using a Cytometers FC500 (Beckman Coulter, Brea, CA) for CD45+ events, representing the numbers of leukocytes present in infected and control corneas.

Statistical analysis. ELISAs and flow cytometry experiments were each performed at least three times. Means of observations were compared by analysis of variance (ANOVA) or Student’s t test, as indicated in each figure legend, using statistical analysis software (SAS Institute Inc., Cary, NC). Statistical significance was set at a P value of ≤0.05.

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References


