Human dipeptidyl peptidase 4 (hDPP4) was recently identified as the receptor for Middle East respiratory syndrome coronavirus (MERS-CoV) infection, suggesting that other mammalian DPP4 orthologs may also support infection. We demonstrate that mouse DPP4 cannot support MERS-CoV infection. However, employing mouse DPP4 as a scaffold, we identified two critical amino acids (A288L and T330R) that regulate species specificity in the mouse. This knowledge can support the rational design of a mouse-adapted MERS-CoV for rapid assessment of therapeutics.
structural differences are anticipated based on distinct differences in susceptibility to MERS infection.

In the absence of obvious structural differences, we hypothesized that specific amino acids contribute to infection of cells that overexpress chDPP4(273–340). To confirm this, we initially mutated five functionally variant surface amino acids in this region of mDPP4 to the corresponding amino acid from hDPP4 using overlap extension PCR to generate various chDPP4 molecules (Fig. 3A). HEK 293T cells were transfected with the indicated chDPP4 constructs followed by infection with rMERS-CoV-red at an MOI of 1. A chDPP4 molecule containing all five mutations (P282T A288L R289I T330R V340I)—shown as “chDPP4 (5 mutations)” in Fig. 3—promoted highly efficient MERS-CoV infection (Fig. 3B), indicating that one or more of these mutations may be sufficient for infection. Previous structural studies of hDPP4 and the MERS RBD suggest that the interaction may require at least two distinct interactions localized to separate structural domains (10). Our T330R and V340I mutants are localized at or near one domain, while mutations comprising P282T, A288L, and R289I are located in the second domain. Accordingly, each group of mutations (T330R V340I and P282T A288L R289I) was independently introduced into mDPP4 and tested for its capacity to support infection of rMERS-CoV-red (Fig. 3B). Although both groups of mutations influence infection to different degrees, neither set recapitulated the levels of infection seen with all five mutations together (Fig. 3B). Further dissection of each group into

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**FIG 1** Mouse DPP4 (mDPP4) does not support MERS-CoV infection. HEK 293T cells were transfected with 3 μg of plasmid expressing human DPP4 (hDPP4) or hDPP4-Venus fusion (A) and mDPP4 or mDPP4-venus fusion (B). At ~20 h posttransfection, cells were infected with rMERS-CoV-red virus at an MOI of 5. Venus fusion proteins were assessed by fluorescence microscopy at 48 h posttransfection. In independent experiments, infection with rMERS-CoV-red virus was assessed for red cells by fluorescence microscopy at ~18 h postinfection. (C) Western blot analysis demonstrates overexpression of mDPP4 and hDPP4. Extracts were prepared at ~48 h posttransfection using AV lysis buffer (3), and samples were heat inactivated for 60 min at 90°C for removal from a biosafety level 3 (BSL3) facility and resolved on an 8% SDS-PAGE gel. Blots were probed with primary goat-anti-DPP4 polyclonal antibody (R&D Systems) at 1:1,000 in 1× Tris-buffered saline–Tween (TBST) or goat anti-actin polyclonal antibody (Santa Cruz) and detected with a secondary rabbit anti-goat–horseradish peroxidase (HRP)-conjugated antibody (Sigma) at 1:10,000 in 1× TBST in 5% milk. (D) Western blot analysis of MERS-CoV S and N proteins. Lysates were collected at ~18 h postinfection and treated as in panel C. Blots were probed with primary mouse polyclonal antiserum at 1:400, raised to S and N proteins as described previously (3), and detected with a secondary goat anti-mouse–HRP (GE Healthcare) at 1:10,000 in 1× TBST in 5% milk.
single mutants revealed that A288L and T330R were partly responsible for the observed infection from each respective group (Fig. 3B). These results are substantiated by Western blot analysis demonstrating detection of the N protein in infected cells expressing either the A288L or T330R mutation but not in cells expressing chDPP4s with the P282T, R289I, or T330R mutation (Fig. 3D). Combination of these two mutations (chDPP4 A288L T330R) recapitulates the level of infection observed for chDPP4 containing all five mutations and approaches what is observed with hDPP4 (Fig. 3B). Quantitation of MERS-CoV-infected red cells (Fig. 3C) and Western blotting of infected cells (Fig. 3D) substantiate the permissibility of chDPP4 A288L T330R, exhibiting nearly a 1.5-log increase in infection compared to mDPP4 (Fig. 3C). Nevertheless, the mutants did not achieve the level of infection observed with hDPP4, indicating that additional amino acids that differ between mice and humans may also contribute to MERS-CoV infection. Regardless of the DPP4 mutation examined, all were efficiently expressed, and intriguingly, all chDPP4 mutants with the T330R mutation exhibit a shift from a doublet to a single lower-molecular-weight band, potentially indicating glycosylation at this site (Fig. 3D). Additionally, the possibility of a restriction factor in rodent cells could be eliminated since mouse and hamster cells ectopically expressing the indicated DPP4s were permissive to MERS-CoV-red infection, with the exception that mDPP4 does not support infection, as observed with HEK 293T cells (Fig. 4). Our results indicate that a successful infection requires a combination of at least two mutations (A288L and T330R), located at distinct structural domains on blades IV and V of the β-propeller of DPP4. These results are in agreement with previous crystal structure data from hDPP4 and the spike RBD, which suggest the hDPP4 equivalents (L294 and R336) are critical residues for binding and infection (9, 10). Our structure prediction model (Fig. 2B) indicates that A288 and L294 and T330 and R336 exhibit positional differences that may govern distinct functional interactions with the MERS-CoV spike protein RBD. Previous phylogenetic analyses indicate that mouse, bat, and ferret
FIG 3  MERS-CoV infection is dependent upon specific amino acids in DPP4.  (A) Vector NTI protein sequence alignment of hDPP4 (top strand) with chDPP4 (middle strand) and mDPP4 (bottom strand) indicating positions of introduced human mutations with red arrows.  (B) HEK 293T cells were transfected with the indicated DPP4 molecule. At 20 h posttransfection, cells were infected with rMERS-CoV-red virus at MOI of 1, and infection was assessed 18 h postinfection by fluorescence microscopy.  (C) In an independent experiment, cells overexpressing the indicated DPP4 constructs were infected with rMERS-CoV-red virus at MOI of 0.1, 0.01, and 0.001 on six-well plates. At 18 h postinfection, cells were scored at the following MOI: no DPP4 and mDPP4, 0.1; chDPP4 P282T A288L R289I T330R V340I [“chDPP4 (5 mutations)”] and chDPP4 A288L T330R, 0.01; and hDPP4, 0.001. Values were normalized to an MOI of 0.1 and expressed as relative infection at 0.1. Human DPP4, chDPP4 P282T A288L R289I T330R V340I, and chDPP4 A288L T330R showed a significant increase in infection over mDPP4 (*, $P < 0.05$, Student’s $t$ test). (D) Western blots demonstrating overexpression of hDPP4, mDPP4, and each chDPP4 molecule, N protein of infected cells, and β-actin as a loading control. Western blots were prepared and probed as described in Fig. 1C and D.

FIG 4  Human and chimeric DPP4 molecules can support MERS-CoV infection in hamster and mouse cells. (A) Baby hamster kidney 21 (BHK-21) cells were electroporated with the indicated DPP4 molecules. At 20 h posttransfection, cells were seeded in 6-well plates and infected with rMERS-CoV-red at an MOI of 2 at 24 h posttransfection. (B) Mouse NIH 3T3 cells were transfected using Nucleofection (according to the Amaza procedure) with the indicated DPP4 molecules. Cells were seeded into 12-well plates and infected with rMERS-CoV-red at an MOI of 4 at 24 h post-Nucleofection. All infections were assessed at 24 h postinfection by fluorescence microscopy.
DPP4s are highly divergent from primate DPP4s (4, 5), despite the capacity of bat DPP4 to serve as a functional receptor for MERS-CoV infection (4). Moreover, swapping the hDPP4 region containing blades IV and V of the β-propeller into the ferret DPP4 resulted in a gain of infection, although the exact residues that mediate this function were not identified (5). Taken together, these data indicate that DPP4 proteins may be structurally conserved across mammalian species. Incorporation of the A288L and T330R modifications in the context of the mDPP4 will facilitate generation of a mouse model through the production of transgenics and can be used to adapt the MERS-CoV to use mDPP4.

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