

Figure 1 Absolute expression of SARS-CoV-2-related genes in different cell types from the mouse small intestinal epithelium.⁷ Data are shown as colour scale of Log2 of mean copies per million (CPM), with circle size indicating fraction of each cell type expressing the gene.

health conditions such as depression, is reported to reduce COVID-19 severity in humans.⁵ The GI tract is a route of SARS-CoV-2;⁶ however, its unknown if EC cells have any specific capacity for infection that would explain the increased 5-HT in patients with COVID-19, or the SSRI treatment efficacy reported. We, therefore, examined (see online supplemental file 1) the transcriptomes of cells lining the gut wall⁷ for expression of genes associated with SARS-CoV-2 infection, with a focus on EE cell subtypes.

Our focus was on gene expression for proteins implicated or known to be involved as COVID-19 receptors for efficient cell entry; ACE2, BSG and NRP1, associated proteins involved in intracellular trafficking and breakdown; TMPRSS2, FURIN and CTSB, and proteins associated with viral protection; LY6E, IFITM1-3 and IFNAR1-2. We identified that the genes encoding for all of these proteins are expressed within the intestinal epithelium (figure 1). Of the known COVID-19 receptors, Ace2 and Bsg genes are highly expressed in all epithelial cell types. However, the more recently identified receptor, NRP1, is expressed exclusively in hormone-producing EE cells at the gene level.





Single-cell gene expression links SARS-CoV-2 infection and gut serotonin

We read with great interest the paper by Ha *et al*¹ demonstrating that circulating levels of serotonin (5-hydroxytryptamine, 5-HT) are increased in COVID-19 and correlate with disease severity and gastrointestinal symptoms such as diarrhoea. Another recent paper by Lin *et al*² in this journal demonstrated that diarrhoea is the most common GI symptom in patients with COVID-19. Almost all 5-HT in our body is produced by enterochromaffin (EC) cells within the epithelium of the GI tract, which constitute approximately half of all enteroendocrine (EE) cells. Gutderived 5-HT modulates gut peristalsis and exacerbates inflammatory responses by acting as a chemotactic molecule for various immune cells and by triggering cytokine release.³ While most gut epithelial cell types are susceptible to SARS-CoV-2 infection, EE cells have the greatest proportion of cells infected at 12 hours after viral exposure.⁴ In addition, the use of selective serotonin reuptake inhibitors (SSRI), normally prescribed to treat mental

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To examine this further, we deter-

mined which subtypes of EE cells

express Nrp1 (figure 2). We focused

on the major EE cell types containing

peptide-1, ghrelin, neurotensin, soma-

tostatin and 5-HT (figure 2A). EC cells

that express tryptophan hydroxylase

1 (Tph1), the rate-limiting enzyme

for non-neuronal 5-HT synthesis, are

the primary cell type in the gut wall

expressing Nrp1, indicating these cells

may be a route of infection and disease

pathogenesis. We then focused solely

on EC cell gene expression using a

second RNA-seq database⁸ and found

that while all COVID-19-related genes

are expressed in EC cells, Nrp1 has the

greatest enrichment of expression of all

these, of approximately 45-fold greater

expression in EC cells than in non-EC

epithelial cells (figure 2B). Subsequent examination of published work iden-

tifies that NRP1 protein expression is highly colocalised in the gastrointes-

tinal wall with cells that express chro-

TMPRSS2 have a >3-fold increase in

SARS-CoV-2 infection compared with

ACE2 and TMPRSS2 alone.¹⁰ Our

data demonstrate that EC cells are

the only gut cell type that expresses

significant levels of these three SARS-

CoV-2-related genes. This, therefore,

provides a link between EC cells and

the increased diarrhoea,² circulating

5-HT,¹ and efficacy of SSRIs⁵ that are

reported in COVID-19. Experiments

investigating SARS-CoV-2 infectivity in the absence of gut-derived 5-HT would

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provide further evidence of this link.

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Cells expressing NRP1, ACE2 and

mogranin-A, a marker of EC cells.⁹

cholecystokinin,

glucagon-like

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