Should children be vaccinated against COVID-19 now?

Brian Li Han Wong , Mary Elizabeth Ramsay, Shamez N Ladhani

The recent authorisation of vaccines against the novel coronavirus, SARS-CoV-2, raises important questions about prioritising the vaccine to those who are most likely to benefit from protection. Unlike influenza vaccines, it is currently not known whether any of the SARS-CoV-2 vaccines can interrupt onward transmission to others. The decision to vaccinate must, therefore, be made on the direct protection offered by the vaccine to those who are vaccinated. The highest burden of severe disease, hospitalisation and death lies among older adults, raising the question as to whether children and young people should be vaccinated during early deployment of any national immunisation programme.

COVID-19 IN CHILDREN
COVID-19 has affected millions of people since SARS-CoV-2 was identified in December 2019. Children account for 1%–3% of diagnosed COVID-19 cases and have less severe disease and better prognosis than adults, with deaths being extremely rare and mainly affecting adolescents and those with significant underlying comorbidities. Early epidemiological studies suggest that children do not contribute much to the spread of SARS-CoV-2 and that younger children may be less likely to get infected or transmit the virus compared with older children or adults. Most children with SARS-CoV-2 infection are either asymptomatic or develop mild and non-specific respiratory symptoms. Some children have, however, developed paediatric inflammatory multisystem syndrome (PIMS-TS; also known as multisystem inflammatory syndrome in children), which shares features with Kawasaki disease and toxic shock syndrome and typically occurs 2 to 4 weeks after SARS-CoV-2 infection.

VACCINATING CHILDREN
The low rates of severe disease and death associated with SARS-CoV-2 infection in children and young people would suggest that they should not be prioritised for vaccination during early vaccine deployment. SARS-CoV-2 vaccine trials have so far rightly focused on adults, and safety data are currently restricted to adults in vaccine trials, with limited data in children. Additionally, large-scale post-implementation surveillance following mass immunisation will be needed to assess the risk of rare events, including antibody-dependent enhancement and PIMS-TS in vaccinated individuals.

HIGH-RISK CHILDREN
Specific paediatric risk groups may benefit from immunisation during the early vaccine deployment stage. Given the limited data on safety and effectiveness of SARS-CoV-2 vaccines in children at present, however, it may be prudent to initially recommend any vaccination for older children (eg, ≥12 years old) who appear to be at risk of severe and fatal disease than younger children and then extend recommendation to children of all ages as more data become available. Since severe COVID-19 is rare in children, however, there are limited data on relative risks of hospitalisation, paediatric intensive care unit (PICU) admission or death in children with specific comorbidities unlike adults. Additionally, children with the highest risk of COVID-19 may have shielded and thus reduced their risk of exposure to the virus. For non-COVID-19 infections, younger age, underlying pulmonary pathology and immunocompromising conditions were associated with more severe outcomes in children.

During the first wave of the pandemic in England, there were eight deaths in SARS-CoV-2 positive children, including four that were attributed to COVID-19, of which three were in children aged 10–15 years with severe underlying neurodisabilities. A large prospective UK study reported neurological conditions (11%; 63/614) as the most prevalent underlying condition in children hospitalised with COVID-19. In a small cross-sectional study of 46 North American PICUs, 24 of 48 (50%) children aged ≤21 years had at least one comorbidity, including 19 (40%) with complex medical problems, defined as children with long-term dependence on technological support (including tracheostomy) associated with developmental delay and/or genetic anomalies. Of note, analysis of a national UK cohort study of adults with COVID-19 identified Down syndrome to be the single most important risk factor for hospitalisation and death (table 1). Adults with cerebral palsy also had a higher risk of hospitalisation and death due to COVID-19, while the risk was only slightly higher in adults with learning disabilities other than Down syndrome. While such data are currently limited in children, it is biologically plausible that Down syndrome and cerebral palsy may also be associated with an increased risk of severe COVID-19.

Considering all the data so far, children with varying degrees and severities of neurodisabilities appear to be over-represented among the small number of cases reported with severe or fatal COVID-19. Given the well-described high risk of large outbreaks in closed institutional settings, one potential group that might benefit from early immunisation could be children with severe neurodisabilities, including cerebral palsy, residing in or in regular attendance at special educational needs or disabilities schools and colleges, paediatric rehabilitation centres, residential schools and care homes for such children. This might give direct protection against COVID-19 in settings where many of the most vulnerable of children might be together. This is also a well-defined group that could easily be identified for targeted immunisation in the early vaccine deployment stage. This same cohort of children with severe and profound learning disability form part of the group with chronic neurological diseases that is recommended for annual influenza vaccination in the UK.

Another potential group for vaccination early in vaccine deployment would be adolescents aged 16–17 years working in health and care settings (especially care homes) who are likely to have similar occupational risks of SARS-CoV-2 exposure as their adult counterparts.

OTHER RISK GROUPS
Although data are limited in children, there are other potential groups that may also be at increased risk of severe and/or fatal COVID-19 such as the immunocompromised, including (stem cell) transplant recipients and haemoglobinopathies such as sickle cell disease as has been reported in adults. Children
with immunocompromising conditions including malignancies represented 8% (48/615) of COVID-19 hospitalisations in the UK and 23% (11/48 children) of PICU admissions for severe COVID-19 in North America.5 6 Vaccine efficacy may be lower in immunocompromised children, so this group may require additional vaccine doses and/or different scheduling compared with immune-competent children. Further vaccine immunogenicity, effectiveness and safety data are needed before recommending vaccination of this highly vulnerable group. Additionally, the current investigational SARS-CoV-2 vaccines have different mechanisms of action, and it is likely some vaccines may be more immunogenic and efficacious for immunocompromised individuals than others.

LONGER TERM CONSIDERATIONS

As more safety and effectiveness data emerge and vaccine availability increases, wider recommendations for vaccination can be considered, particularly for other high-risk children. It is likely that the major risk groups for influenza vaccination may also be applicable to SARS-CoV-2 and therefore, over time, these same risk groups—especially those with chronic respiratory conditions including ex-premature infants with chronic lung disease and those with chronic heart disease—could also be recommended SARS-CoV-2 vaccination.3 7 Whether all children will eventually be vaccinated will depend on a number of factors. These include the role of children in transmission as well as the safety, effectiveness and duration of protection afforded by available vaccines. Most importantly, it will depend on whether such vaccines interrupt transmission, thereby inducing indirect (herd) protection for the population, as has been demonstrated with the UK childhood influenza immunisation programme. Up-to-date recommendations for SARS-CoV-2 vaccination in the UK will be maintained within the ‘Green Book’ (Immunisation Against Infectious Disease).7

Table 1  Adjusted HR (95% CI) of death from COVID-19 in men and women, adjusted for age, ethnicity, deprivation, body mass index and a range of comorbidities (adapted from Clift et al)3

<table>
<thead>
<tr>
<th>Condition</th>
<th>Men</th>
<th>Women</th>
</tr>
</thead>
<tbody>
<tr>
<td>Down syndrome</td>
<td>9.80 (4.62 to 20.78)</td>
<td>32.55 (18.13 to 58.42)</td>
</tr>
<tr>
<td>Bone marrow or stem cell transplant in past 6 months</td>
<td>6.10 (1.11 to 33.54)</td>
<td>2.78 (0.22 to 34.55)</td>
</tr>
<tr>
<td>Sickle cell disease or severe immunodeficiency</td>
<td>4.41 (1.41 to 13.81)</td>
<td>5.94 (1.89 to 18.67)</td>
</tr>
<tr>
<td>Cerebral palsy</td>
<td>3.45 (1.10 to 10.78)</td>
<td>3.45 (1.10 to 10.78)</td>
</tr>
<tr>
<td>Learning disability apart from Down syndrome</td>
<td>1.36 (1.14 to 1.60)</td>
<td>1.36 (1.11 to 1.65)</td>
</tr>
</tbody>
</table>

Correction notice This article has been corrected since it first published. The provenance and peer review statement has been included.

Twitter Brian Li Han Wong @brianwong_

Contributors All authors contributed equally to the manuscript.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; internally peer reviewed. This article is made freely available for use in accordance with BMJ’s website terms and conditions for the duration of the covid-19 pandemic or until otherwise determined by BMJ. You may use, download and print the article for any lawful, non-commercial purpose (including text and data mining) provided that all copyright notices and trade marks are retained.

© Author(s) (or their employer(s)) 2021. No commercial re-use. See rights and permissions. Published by BMJ.

To cite Wong BLH, Ramsay ME, Ladhani SN. Arch Dis Child. 2021;105:376-382. doi:10.1136/archdischild-2020-321225

Received 19 November 2020

Revised 10 December 2020

Accepted 11 December 2020

Arch Dis Child 2021;105:376-382. doi:10.1136/archdischild-2020-321225

ORCID iDs

Brian Li Han Wong http://orcid.org/0000-0001-8709-5847

Shamez N Ladhani http://orcid.org/0000-0002-0856-2476

REFERENCES


