UKHSA publishes a weekly national influenza and COVID-19 surveillance report which summaries the information from the surveillance systems which are used to monitor influenza, COVID-19 and other seasonal respiratory viruses in England.

Additional figures based on these surveillance systems are included in this slide set.

The figures presented in this slide set are based on data from week 40 (between 3 October and 9 October 2022).
Contents

1) COVID-19 Pandemic Overview
2) Confirmed COVID-19 episodes in England
3) Respiratory Datamart system (England)
4) Second generation surveillance system (SGSS)
5) Community surveillance
6) Surveillance in ‘educational-age’ cohorts
7) Secondary Care surveillance
8) Mortality surveillance
9) Possible reinfections in England
10) Co/secondary infections with COVID-19
COVID-19 Pandemic Overview
Confirmed COVID-19 episodes tested under Pillar 1, by sample week, since week 5 2020
Confirmed COVID-19 episodes tested under Pillar 2, by sample week, since week 5 2020
Weekly overall hospital and ICU/HDU admission rates per 100,000 of new COVID-19 cases reported through SARI Watch, England since week 12 2020.
Since 1 April 2022, free universal symptomatic and asymptomatic testing for the general public in England is no longer available, as outlined in the plan for living with COVID-19. Data should be interpreted in the context of this change to testing.
Confirmed COVID-19 episodes in England
Confirmed COVID-19 episodes in England

Data Information

• From the week 32 report onwards, case rates have been updated to use the latest ONS population estimates for mid-2020. Previously case rates were calculated using the mid-2019 population estimates.

• From 11 January 2022 the requirement for confirmatory PCR testing in individuals who test positive using a lateral flow device was temporarily removed.

• Rates by ethnicity and IMD quantile will continue to be presented using the mid-2019 estimates, until the mid-2020 estimates become available.

• From 31 January 2022, UKHSA moved all COVID-19 case reporting in England to use a new episode-based definition which includes possible reinfections. Each infection episode is counted separately if there are at least 91 days between positive test results (PCR or LFD). Each infection episode begins with the earliest positive specimen date. Further information can be found on the UK COVID-19 dashboard.

• Since 1 April 2022, free universal symptomatic and asymptomatic testing for the general public in England is no longer available, as outlined in the plan for living with COVID-19. As such, there will be a reduction in the reporting of data obtained through Pillar 2 from April 2022 onwards. Data in this report should be interpreted in the context of this change to testing. Public health guidance remains in place for cases and their close contacts.
Confirmed COVID-19 episodes tested under Pillar 2, based on sample week with overall weekly PCR positivity for Pillar 2 (%)
Weekly confirmed COVID-19 case rates per 100,000, by episode, tested under Pillar 2, by sex
Weekly confirmed COVID-19 case rates per 100,000, by episode, tested under Pillar 2, by age group
Weekly PCR positivity (%) of confirmed COVID-19 cases tested overall and by sex under Pillar 2
Weekly PCR positivity (%) of confirmed COVID-19 cases tested under Pillar 2, by male and age group
Weekly PCR positivity (%) of confirmed COVID-19 cases tested under Pillar 2, by female and age group.
Weekly confirmed COVID-19 case rates by episode, per 100,000 population (Pillar 2), by UKHSA centres and sample week.
Weekly PCR positivity of confirmed COVID-19 cases tested under Pillar 2 (%) by UKHSA centres and sample week
Weekly incidence per 100,000 population by ethnicity (Pillar 2), England
Weekly COVID-19 episodes tested under Pillar 1, per 100,000 population by age group and region, weeks 31 to 40
Weekly COVID-19 episodes tested under Pillar 2, per 100,000 population by age group and region, weeks 31 to 40
Weekly COVID-19 episodes tested under Pillar 1, per 100,000 population by ethnicity and region, weeks 31 to 40.
Weekly COVID-19 episodes tested under Pillar 2 per 100,000 population by ethnicity and region, weeks 31 to 40
Weekly COVID-19 rate tested under Pillar 1, per 100,000 population by IMD quintile (1 being the most deprived and 5 being the least deprived)

![Graph showing weekly COVID-19 rates tested under Pillar 1, per 100,000 population by IMD quintile.]
Weekly COVID-19 rate tested under Pillar 2, per 100,000 population by IMD quintile (1 being the most deprived and 5 being the least deprived)
Weekly rate of COVID-19 episodes per 100,000 population (Pillar 2), by upper-tier local authority, England (box shows enlarged map of London area)
Cumulative rate of COVID-19 episodes per 100,000 population tested under Pillar 1, by upper-tier local authority, England (box shows enlarged map of London area).
Cumulative rate of COVID-19 episodes per 100,000 population tested under Pillar 2, by upper-tier local authority, England (box shows enlarged map of London area)
Respiratory Datamart system (England)
Respiratory DataMart – Respiratory syncytial virus (RSV)
Respiratory DataMart – Respiratory syncytial virus (RSV) weekly positivity by UKHSA region

Proportion positive (%) vs Week number for:
- North of England
- South of England
- London
- Midlands and East of England
Respiratory DataMart – other respiratory viruses

**Adenovirus**

- Number of positive samples
- Proportion positive (%)

**Parainfluenza**

- Number of positive samples
- Proportion positive (%)

**Rhinovirus**

- Number of positive samples
- Proportion positive (%)

**hMPV**

- Number of positive samples
- Proportion positive (%)

13 October 2022
Second generation surveillance system (SGSS)
The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.

Previously, this data was presented by report date however is now presented by specimen date.
SGSS reported Influenza B cases by region (all ages)

The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.

Previously, this data was presented by report date however is now presented by specimen date.
SGSS reported RSV cases by region (all ages)

The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution. Previously, this data was presented by report date however is now presented by specimen date.
The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.
SGSS reported Parainfluenza cases by region (all ages)

The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.
SGSS reported Rhinovirus cases by region (all ages)

The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.
The presented figures are based on laboratory reports through SGSS. Testing and reporting procedures vary by virus, UKHSA centre and over time, including short-term trends in testing. Therefore comparisons should be done with caution.
Community surveillance
Acute respiratory infection (ARI) outbreaks linked to educational settings

**Data Information**
We report on new acute respiratory infection (ARI) incidents reported to Health Protection Teams (HPTs) and entered on HPZone in the previous reporting week in educational settings by locality. The incidents captured on HPZone represent a subset of all ongoing clusters and outbreaks in England. A variety of arrangements are in place with local authorities and other stakeholders supporting HPTs, however data may not routinely be documented on HPZone. As a result, the number of outbreaks reported for some of the regions are underestimates.
Number of acute respiratory infection outbreaks reported to UKHSA by type of educational setting, England
Number of acute respiratory infection outbreaks by type of educational setting, England

End of academic year total
Week 35 2021-34 2022

<table>
<thead>
<tr>
<th>PHE Centres</th>
<th>Nursery</th>
<th>Primary School</th>
<th>Secondary School</th>
<th>Combined</th>
<th>Special Educational Needs (SEN) schools</th>
<th>College University</th>
<th>Total</th>
</tr>
</thead>
<tbody>
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<td>Total</td>
<td>540</td>
<td>1761</td>
<td>596</td>
<td>161</td>
<td>1306</td>
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Week 40 2022
Main table

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<th>PHE Centres</th>
<th>Nursery</th>
<th>Primary School</th>
<th>Secondary School</th>
<th>Combined</th>
<th>Special Educational Needs (SEN) schools</th>
<th>College University</th>
<th>Total</th>
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<td>1 (0)</td>
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<td>North East Centre</td>
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<td>0 (0)</td>
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<tr>
<td>North West Center</td>
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<td>South East Centre</td>
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<tr>
<td>South West Centre</td>
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<td>0 (0)</td>
<td>3 (0)</td>
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<td>4 (0)</td>
</tr>
<tr>
<td>West Midlands Centre</td>
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<td>0 (0)</td>
<td>2 (1)</td>
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<tr>
<td>Yorkshire &amp; the Humber</td>
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<td>0 (0)</td>
<td>10 (4)</td>
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<td>20 (6)</td>
</tr>
</tbody>
</table>

* Number of acute respiratory infection for the most recent week in brackets
Secondary Care surveillance
Weekly admission rates for hospital and ICU/HDU laboratory confirmed COVID-19 cases reported through SARI Watch, week 40

Source: UKHSA SARI-Watch (Severe Acute Respiratory Infection-Watch, formerly CHESS).

*Only NHS Acute trusts that have reported ≥1 day in the past week; excludes Specialist trusts. Acute NHS trusts (including Specialist trusts) reporting into SARI-Watch COVID-19 hospitalisation surveillance are typically around 100 per week. This was 87 for the hospitalisation (all levels of care) indicator in week 3 October 2022 to 9 October inclusive and 81 trusts for the ICU/HDU indicator. For the maps, as Specialist trusts are excluded, the number of trusts providing data on COVID-19 hospitalisations in week ending 9 October 2022 was 78 and 74 for ICU/HDU admissions for COVID-19.
Age/sex pyramid of hospitalisations (all levels of care) for COVID-19, data from sentinel acute NHS trusts, England

(a) Peak of 2\textsuperscript{nd} wave (week 53 2020 to week 3 2021) n= 6,359

Reporting trusts=22

(b) Most recent 4 weeks (week 37 2022 to 40 2022) n= 809

Reporting trusts=10
Age/sex pyramid for admissions to ICU/HDU for COVID-19, mandatory case level data, acute NHS trusts, England

(a) Peak of 2\textsuperscript{nd} wave (week 53 2020 to week 3 2021) n= 3,349

(b) Most recent 4 weeks (week 37 2022 to 40 2022) n=65

Reporting trusts=70

Reporting trusts= 25
Laboratory confirmed admissions for COVID-19, to acute NHS trusts, by level of care and ethnicity

(a) Peak of 2nd wave (week 53 2020 to week 3 2021)

(b) Most recent 4 weeks (week 37 2022 to 40 2022)

Reporting trusts
Lower level of care=21
ICU/HDU=68

Reporting trusts
Lower level of care=10
ICU/HDU=23
Weekly COVID-19 hospitalisation rate per 100,000 trust catchment population by age group and region, weeks 31 to 40.
Caveat: From week 24 (2021) the ethnicity analysis is based on a new method for assigning ethnicity, developed by UKHSA. The previous method used the most recent ethnicity recorded through linkage to Hospital Episode Statistics. However, this method led to unfeasibly high rates in the ‘Other’ ethnic group when applied to COVID-19 cases, hospitalisation or mortality. The new method uses the most frequent ethnicity recorded through linkage to Hospital Episode Statistics, unless the most frequent was ‘Other’ when the second most frequent was chosen.
Rate of admission to ICU/HDU by ethnicity, per 100,000 trust catchment population

Caveat: From week (24 2021) the ethnicity analysis is based on a new method for assigning ethnicity, developed by UKHSA. The previous method used the most recent ethnicity recorded through linkage to Hospital Episode Statistics. However, this method led to unfeasibly high rates in the ‘Other’ ethnic group when applied to COVID-19 cases, hospitalisation or mortality. The new method uses the most frequent ethnicity recorded through linkage to Hospital Episode Statistics, unless the most frequent was ‘Other’ when the second most frequent was chosen.
Notes
1) Case-level sentinel data from SARI-Watch, form week 35 2021 (commencing 30 August 2021) to week 39 2022 (ending 2 October 2022) inclusive
2) Total 28,452 records in period of analysis, of which 34% (n=9,727) had COVID-19 as primary reason for admission ('Yes').
3) SARS-CoV-2 patients with evidence of COVID-19 treatment but have 'No' or 'Unknown' for COVID-19 as primary reason for admission (n=960) are reassigned to COVID-19 as primary reason of admission ('Yes').
4) Reassignment increases COVID-19 as primary reason for admission ('Yes') from 9,727 to 10,687
5) 24% (6,804/28,452) of total records in this period have missing data on the 'Admission due to COVID-19' indicator – these are excluded from analysis
6) Caveats: London trusts under-represented and most recent weeks are subject to retrospective updates
SARS-CoV-2 Whole Genome Sequencing (WGS) coverage, England
Coverage of sequencing with a valid result and genotyping over time (3 October 2021 to 3 October 2022)

Coverage of sequencing with a valid result and genotyping over time (3 October 2021 to 3 October 2022).

Episodes where the individual only tested using a lateral flow device are excluded. Grey shading was applied to the previous 14 days to account for reporting delays in sequencing data.
Preceding/co-/secondary infections with COVID-19

Slides for weekly covid flu report
Preceding/co-/secondary infections with COVID-19

- Caveat - undertesting for other pathogens may result in an underestimate of preceding/co-/secondary infection cases.

- Preceding/co-/secondary infections refers to when a patient has a COVID-19 infection with one or more other pathogen (Please see Appendix 1 – Pre-/co-/secondary infection with COVID-19 definitions.)
  - Preceding infection: SARS-CoV-2 acquired after another pathogen
  - Co-infection: SARS-CoV-2 and other pathogen acquired at the same time
  - Secondary infection: SARS-CoV-2 acquired before another pathogen

- Numbers of preceding/co-/secondary infection remain low across UKHSA surveillance systems.

- For patients with severe respiratory failure requiring Extra Corporeal Membrane Oxygenation (ECMO), analysis of data from five adult ECMO centres in England indicates that among patients with severe respiratory failure due to COVID-19, almost a third of these have co/secondary infections.

- Published data analysis from pandemic wave 1 (W-1) indicates increased mortality associated with COVID-19 and influenza, key bacterial and fungal infections and invasive pneumococcal disease (IPD) in comparison to patients without co/secondary infection.

- Data analysis from W-1 indicates that Aspergillus and candidemia cases have increased risk of mortality in comparison to patients without co/secondary infection.
Co/secondary infections among patients with severe respiratory failure requiring Extra Corporeal Membrane Oxygenation (ECMO) 

Analysis is based on cumulative data from five adult ECMO centres in England. Surveillance is all year round. Each season commences around October (ISO week 40) ending in September (ISO week 39) in the following year.

Data for the current season (2021-22) is from 4 October 2021 to 2 October 2022 inclusive (week 40 2021 to week 39 2022). This period includes effects from the Delta and Omicron waves of the pandemic. The 2020-21 season is from 28 September 2020 to 3 October 2021 inclusive and includes effects from the Alpha and Delta waves. The 2019-20 season is from 30 September 2019 to 27 September 2020 inclusive and includes effects from the original Wuhan strain.

- In the 2021-22 season, 40% (31/77) of ECMO patients admitted for severe respiratory failure due to laboratory confirmed COVID-19 had clinically significant co/secondary infections. In the previous season (2020-21) this proportion was 33% (134/402). In the 2019-20 season this proportion was 33% (79/236).

- In all three seasons the majority of clinically significant co/secondary infections among respiratory failure COVID-19 cases comprised Gram-negative bacilli from the order Enterobacterales:
  - 45% (14/31) in the current season 2021-22
  - The decrease in 2020-21 compared to 2019-20 reached borderline significance (p=0.057). No change was detected in other key pathogens between these two time periods.
  - No evidence of change was detected in the proportion of co/secondary infections due to Enterobacterales between 2020-21 and 2021-2022.

HCAI, Fungal, AMR, AMU & Sepsis Division
Updates

- The following outputs included in this section have been produced via the Unified Infection Dataset (UID), combining previously separate data pipelines
  - Key HCAI bacterial and fungal specimens reported to SGSS and HCAI data capture system
  - Respiratory viral specimens reported to SGSS and Respiratory Datamart
  - Fungal specimens reported to mycology reference lab (MRL)
- Data are provisional and subject to change due to possible delayed reporting of microbiological samples
- Free community testing ended 31 March 2022 as part of the government’s Living with COVID-19 plan, with asymptomatic testing continuing in some settings. As of 31 August 2022, asymptomatic testing in all settings, including hospitals, has been paused. Please use caution when comparing incidence of bacterial, fungal and viral preceding/co-/secondary infections over time due to these differences in testing strategies.
Number of COVID-19 patient-episodes with bacterial, fungal or respiratory viral infections in COVID-19 patients diagnosed in England during wave 3*, by infection type and timing of diagnosis

![Table](image)

* SARS-CoV2 specimen dates from 27 Apr 2021 to 4 Sep 2022 (N=15,913,127). Last updated 7 Oct 2022.

**Definition for secondary infection differs for MRL specimens - detection within 60 days**

† includes the combination Bacterial/fungal bloodstream & *Clostridioides difficile* infection (12 preceding, 1 coinfection & 21 secondary), Bacterial/fungal bloodstream, lower respiratory & *Clostridioides difficile* infection (1 secondary), & Bacterial/fungal lower respiratory & *Clostridioides difficile* infection (1 preceding & 3 secondary)

**Key findings:**

- 0.1% of COVID-19 patient-episodes had a bacterial, fungal or other respiratory viral infection detected in either the 28 days prior or following (60 days following for MRL) their COVID-19 diagnosis.
- Prevalence in W3 lower than W2 and W1; however, patient-episodes of COVID-19 and another key infections are still higher in W3 than W2 and W1 (19,850 vs 11,830 vs 4,636, respectively).
- Most infections with key organisms were categorised as secondary infections (39.7%).

Please see appendix 1 for pre-/co-/secondary infection definitions with SARS-CoV-2

Please note patients can have multiple COVID-19 infection-episodes, numbers here do not reflect the number of patients.
Most frequent bacterial/fungal species in blood or lower respiratory tract specimens, by timing of diagnosis, in COVID-19 patients diagnosed in England during wave 3

Key findings:
In wave 3, the most frequent bacterial/fungal organisms identified from blood specimens were *Escherichia coli*, *Staphylococcus aureus* and *Klebsiella pneumoniae* and from respiratory specimens were *S. aureus*, *Pseudomonas aeruginosa* and *E. coli*.}

Please note different y-axes
Most frequent viral specimens, by timing of diagnosis, in COVID-19 patients diagnosed in England during wave 3

Key findings:
In wave 3, the most frequent viral organisms identified from respiratory specimens were RSV, rhinovirus and influenza A.
Most frequent fungal species (MRL), by timing of diagnosis, in COVID-19 patients diagnosed in England during wave 3

Key findings:
In wave 3, the most frequent fungal organisms identified were Aspergillus fumigatus complex and Candida albicans.
## Appendix 1: Pre-/co-/secondary infection definitions with COVID-19

The day pertains to the date of the sample collection that yielded a positive result. These definitions do not apply to persistent COVID-19 patients. Patients with persistent COVID-19 require independent clinical assessment.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Definition co-infection with SARS-CoV-2 †</th>
<th>Definition of infection pre-SARS-CoV-2 infection (other pathogen is primary infection) or Definition of post SARS-CoV-2 secondary infection (SARS-CoV-2 is primary infection)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influenza A</td>
<td>+/- 1d</td>
<td>2-28d^</td>
</tr>
<tr>
<td>Influenza B</td>
<td>+/- 1d</td>
<td>2-28d^</td>
</tr>
<tr>
<td>RSV</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Adenovirus</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Enterovirus</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Human metapneumovirus</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Parainfluenza (any subtype)</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Seasonal coronavirus</td>
<td>+/- 1d *</td>
<td>2-28d</td>
</tr>
<tr>
<td>Rhinovirus</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Co-infections in ECMO patient (patients with most severe clinical respiratory signs)</td>
<td>Individual case review</td>
<td>Individual case review</td>
</tr>
<tr>
<td>ECMO patients</td>
<td>Individual case review</td>
<td></td>
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<tr>
<td>Blood stream and respiratory infections (bacterial and fungal)</td>
<td>Individual case review</td>
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<tr>
<td>Achromobacter xylosoxidans</td>
<td>+/- 1d</td>
<td>2-28d</td>
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<td>Acinetobacter spp.</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Aspergillus</td>
<td>+/- 1d</td>
<td>2-28d (pre) 2-60d (post, continually hospitalised patients only)</td>
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<tr>
<td>Bordetella pertussis</td>
<td>+/- 28 d Culture/PCR (based on pertussis sample date)</td>
<td>N/A (Pertussis presentation is often delayed)</td>
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<td></td>
<td>+/- 28 Serology/Oral fluid (anti-pertussis toxin Ig)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(based on pertussis symptom onset date, excluding cases without onset date)</td>
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<td>Burkholderia cepacia</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Candida spp</td>
<td>+/- 1d</td>
<td>2-28d (pre) 2-60d (post, continually hospitalised patients only)</td>
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<tr>
<td>Chlamydia pneumoniae</td>
<td>0-7d PCR</td>
<td>PCR within 14-28 d (8-13d PCR^)</td>
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<td>Enterobacter spp.</td>
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<td>Enterococcus spp.</td>
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<td>2-28d</td>
</tr>
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<td>E. coli</td>
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<td>2-28d</td>
</tr>
<tr>
<td>Haemophilus influenzae</td>
<td>+/- 2d</td>
<td>3-28d</td>
</tr>
</tbody>
</table>

Continued overleaf
Appendix 1 continued: Pre-/co-/secondary infection definitions with COVID-19

The day pertains to the date of the sample collection that yielded a positive result. These definitions do not apply to persistent COVID-19 patients. Patients with persistent COVID-19 require independent clinical assessment.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Definition co-infection with SARS-CoV-2 †</th>
<th>Definition of infection pre-SARS-CoV-2 infection (other pathogen is primary infection) or Definition of post SARS-CoV-2 secondary infection (SARS-CoV-2 is primary infection)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood stream and respiratory infections (bacterial and fungal)</td>
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<tr>
<td>Klebsiella spp.</td>
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<tr>
<td>Legionella pneumophilia/species</td>
<td>Individual case review</td>
<td>Individual case review</td>
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<tr>
<td>Mycoplasma pneumoniae</td>
<td>0-7d PCR, IgM serology 0-21d &lt;16y</td>
<td>PCR within 14-28 d (8-13d PCR*)</td>
</tr>
<tr>
<td>Neisseria meningitidis</td>
<td>+/- 2d</td>
<td>3-28d</td>
</tr>
<tr>
<td>Pseudomonas spp.</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Serratia spp.</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>+/- 1d</td>
<td>2-28d</td>
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<td>Coag-neg Staphylococcus (S. haemolyticus)</td>
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<td>Streptococcus spp. ‡</td>
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<td>Streptococcus pneumoniae</td>
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<td>Mycobacterium tuberculosis</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>HIV</td>
<td>Individual case review</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Gastrointestinal infections</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Listeria</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Campylobacter</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Shiga toxin-producing E. coli (STEC)</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Norovirus</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Salmonella</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Shigella</td>
<td>0-5d *</td>
<td>Individual case review</td>
</tr>
<tr>
<td>Anaerobes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C. difficile</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
<tr>
<td>Bacteroides sp. (B. fragilis and non-fragilis Bacteroides)</td>
<td>+/- 1d</td>
<td>2-28d</td>
</tr>
</tbody>
</table>

See next slides for notes
Appendix 1 continued: Pre-/co-/secondary infection definitions with COVID-19

**Notes**
† From the first specimen date of a SARS-CoV-2 patient episode.
* Additional data check required. (Resistance is not detailed, data for MERS is not currently available).
^ Definition post- SARS-CoV-2 secondary infection (SARS-CoV-2 is primary infection). This has been extended from prior 14d secondary infection definition for influenza used by UKHSA to account for disparities in testing throughout the 28d period after SARS-CoV-2 detection.
‡ Streptococcus species includes the following groups and species:

<table>
<thead>
<tr>
<th>Group</th>
<th>Species/other names</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anginosus Group</td>
<td>Streptococcus anginosus; Streptococcus constellatus (Streptococcus constellatus subspecies pharynges); Streptococcus Group F; Streptococcus intermedius; Streptococcus milleri group; Streptococcus sinensis</td>
</tr>
<tr>
<td>Bovis Group</td>
<td>Streptococcus alactolyticus; Streptococcus bovis untyped; Streptococcus equinus; Streptococcus galloyticus subspecies galloyticus (Streptococcus bovis biotype I); Streptococcus infantarius (Streptococcus infantarius sp infantarius; Streptococcus bovis biotype II); Streptococcus lutetiensis; Streptococcus infantarius subspecies coli (Streptococcus bovis biotype II); Streptococcus pasteurianus (Streptococcus bovis biotype II)</td>
</tr>
<tr>
<td>Closely Related Genera</td>
<td>Abiotrophia spp.; Aerococcus spp.; Faklamia spp.; Gemella spp.; Globicatella sanguinis; Granulicatella spp.; Leuconostoc spp.; Pedicoccus spp.; Peptostreptococcus spp.</td>
</tr>
<tr>
<td>Mutans Group</td>
<td>Streptococcus mutans; Streptococcus sobrinus</td>
</tr>
<tr>
<td>Other streptococci (including but not limited to)</td>
<td>Anaerobic streptococcus; Streptococcus acidominimus; Streptococcus spp., other named/not fully identified; Streptococcus suis; Streptococcus uberis</td>
</tr>
<tr>
<td>Salivarius Group</td>
<td>Streptococcus vestibularis; Streptococcus thermophilus</td>
</tr>
<tr>
<td>Sanguinis Group</td>
<td>Streptococcus gordonii; Streptococcus massiliensis; Streptococcus parasanguinis; Streptococcus sanguinis</td>
</tr>
<tr>
<td>Streptococcus Group A</td>
<td>Group A; Streptococcus pyogenes; Streptococcus dysgalactiae subspecies equisimilis</td>
</tr>
<tr>
<td>Streptococcus Group B</td>
<td>Group B; Streptococcus agalactiae</td>
</tr>
<tr>
<td>Streptococcus Group C</td>
<td>Group C; Streptococcus dysgalactiae subspecies equisimilis; Streptococcus equi subspecies zooepidemicus</td>
</tr>
<tr>
<td>Streptococcus Group G</td>
<td>Group G; Streptococcus canis; Streptococcus dysgalactiae subspecies equisimilis</td>
</tr>
</tbody>
</table>