# MinION Mk1B IT requirements

#### **Overview**

The MinION $^{\text{m}}$  is a small, self-contained device for nanopore sequencing. It plugs directly into a USB Type-A port or a USB Type-C port with a recommended adapter (5 Gbps).

The MinION device is controlled by the MinKNOW™ software.

MinKNOW carries out several core tasks. These include data acquisition, real-time analysis and feedback, basecalling, data streaming, device control including selecting the run parameters, sample identification and tracking. Furthermore, it ensures that the platform chemistry is performing correctly to run the samples.

Despite its small size, the MinION Mk1B can sequence many gigabases (Gbases) in a single sequencing experiment. As such, it is recommended that users purchase a powerful computer to ensure all the technology features are accessible.

The default data analysis workflow when starting to use the MinION is below:



## **Host computer specification**

From MinKNOW v23.07 onwards, our new basecaller Dorado is integrated into MinKNOW. This enables accelerated basecalling on both NVIDIA GPUs and Apple Silicon. We strongly recommend using an NVIDIA GPU or Apple Silicon Mac if you want basecalling to keep up with the rate of data generation.

| Component        | Required specification: GPU high-accuracy basecalling | Required specification:<br>data acquisition/CPU<br>basecalling<br>(note: CPU basecalling<br>performance is limited<br>- a GPU is<br>recommended) |
|------------------|---|--|
| Operating system | Windows - 10<br>Linux - Ubuntu 20.04 and 18.04        | Windows - 10<br>macOS - Mojave, Catalina<br>Linux - Ubuntu 20.04 and<br>18.04  |
| Memory/RAM       | 16 GB RAM or higher                                   | 16 GB RAM or higher  |

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|-----------|--|--|
| CPU       | Intel i7, i9, Xeon, or better, with at least 4 cores/8 threads  Ryzen 5, 7, or better, with at least 4 cores/8 threads   | Intel i7, i9, Xeon, or better, with at least 4 cores/8 threads  Ryzen 5, 7, or better, with at least 4 cores/8 threads                           |
| GPU       | NVIDIA GPU RTX 2060 SUPER or better, with at least 8 GB of GPU memory. Technical information can be found on various websites, for example https://www.techpowerup.com/gpu-specs/.  Widely-available examples include the RTX 2060 SUPER, RTX 2070, RTX 3060, RTX 3070. Ampere-based GPUs (the 3000 series, A series etc.) are particularly recommended for optimal performance.  If you are working with a different type of GPU than the models listed above, please ensure that it has a CUDA Compute Capability >6.1 (for more information about CUDA-enabled GPUs, see the NVIDIA website). |  |
| Storage   | 1 TB internal SSD or higher  | 1 TB internal SSD or higher  |
| Ports     | USB3.0   | USB3.0   |

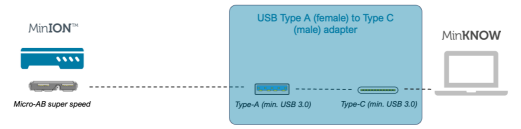
We recommend internal solid-state storage for MinKNOW installation as well as data output/acquisition. Solid-state drives are much faster than traditional hard drives and are able to keep up with the flow of data generated during a sequencing run.

#### **Computer recommendations**

Below are some examples of models/workstations that meet or can be configured to meet the recommended specification for the MinION Mk1B. We have not extensively tested any of these computers, so purchasing is at your own risk. Ensure that you have configured the computer to meet the specification in the table above. Other manufacturers are available; check the availability for your region.

| Performance band                          | Example laptop                            |  |
|---|---|--|
| Low: Fast basecalling model keep-up       | M1 MacBook Air (2023)                     |  |
| Medium: High accuracy basecalling keep-up | M2 Max 16" MacBook Pro (2023) / MacStudio |  |

Connecting the MinION Mk1B to a computer without a USB Type-A port



If your laptop does not have a USB Type-A port, we recommend using a USB Type-A to USB Type-C adapter (Female to Male 5 Gbps). Below is a non-exhaustive list of example adapters. We have not extensively tested all models below, so purchasing is at your own risk

- StarTech 6"
- RS Pro 150 mm
- Belkin

The below requirements are to ensure correct operation of the MinION during a sequencing experiment:

| Component                             | Minimum requirement  |
|---------------------------------------|--|
| User<br>account<br>privilege<br>level | Local Administrator privileges are needed for MinKNOW installation and updates.  Local Administrator privileges are not required for running sequencing experiments.   |
| Internet<br>connection                | An internet connection is required at all times for software updates and telemetry.  Offline configurations can be made available for field use and expeditions. Please contact support@nanoporetech.com if you are planning to use your MinION offline.   |
| Antivirus<br>settings                 | Antivirus software scanning the entire hard drive uses significant amounts of resources and may cause performance issues. For this reason, it is recommended that the antivirus scans are switched to manual setting, rather than to automatic, so that they can be performed when the MinION is not in use. |
| OS update settings                    | OS system updates should be switched to manual mode, as downloading them during a sequencing experiment may impact performance.  Updates that require restart will fatally halt a run.   |
| Telemetry<br>feedback                 | HTTPS/port 443 to 52.17.110.146, 52.31.111.95, 79.125.100.3 (outbound-only access) or DNS rule for ping.oxfordnanoportal.com   |
| EPI2ME<br>analysis                    | Ethernet: HTTPS/port: 443 TCP access to AWS eu-west-1 IP ranges: http://docs.aws.amazon.com/general/latest/gr/aws-ip-ranges.html   |
| Software updates                      | HTTPS/port 443 to 178.79.175.200 and 96.126.99.215 (outbound-only access) or DNS rule for cdn.oxfordnanoportal.com   |

## **Telemetry**

MinKNOW collects telemetry information during sequencing runs as per the Terms and Conditions to allow monitoring of device performance and enable remote troubleshooting. Some of this information comes from free-form text entry fields, therefore no personally-identifiable information should be included. We do not collect any sequence data.

The EPI2ME platform is hosted within AWS and provides cloud-based analysis solutions for multiple applications. Users upload

sequence data in FASTQ format via the EPI2ME Agent, which processes the data through defined pipelines within the EPI2ME Portal. Downloads from EPI2ME are either in Data+Telemetry or Telemetry form. The EPI2ME portal uses telemetry information to populate reports.

### **Software updates**

Depending on your geographical region, only one of 178.79.175.200 or 96.126.99.215 will be used for provision of updates to device software. The Updates are triggered as pull requests, therefore outbound-only access is required.

### File types

Nanopore sequencing data is stored in three file types: POD5, FASTQ and BAM. Basecalling summary information is stored in a sequencing summary.txt file:

- POD5 is an Oxford Nanopore-developed file format which stores nanopore data in an accessible way and replaces the legacy .fast5 format. This output also reads and writes data faster, uses less compute and has smaller raw data file size than .fast5.
- .fast5 is a legacy file format based upon the .hdf5 file type, which contains all information needed for analysing nanopore sequencing data and tracking it back to its source. A .fast5 file contains data from multiple reads (4000 reads as default), and is several hundred Mb in size.
- FASTQ is a text-based sequence storage format, containing both the sequence of DNA/RNA and its quality scores. By default, nanopore sequencing experiments save up to 4000 DNA sequences in one FASTQ file. File size can vary from <1 Mb to tens of Mb depending on the number and length of sequences. Retaining only FASTQ files will allow use of standard downstream analysis tools using the DNA/RNA sequence, but no further sequence data can be generated when improvements in basecalling become available.
- BAM files are output if you perform alignment or modified base calling on the basecalled dataset.
- sequencing\_summary.txt contains metadata about all basecalled reads from an individual run. Information includes read ID, sequence length, per-read q-score, duration etc. The size of a sequence summary file will depend on the number of reads sequenced.

Example file sizes below are based on different throughputs from an individual flow cell, with a run saving POD5, FASTQ, and BAM files with a read N50 of 23 kb.

| Flow cell output (Gbases) | POD5 storage<br>(Gbytes) | FASTQ.gz storage<br>(Gbytes) | Unaligned BAM with modifications (Gbytes) |
|---------------------------|--------------------------|------------------------------|---|
| 10                        | 70                       | 6.5                          | 6   |
| 15                        | 105                      | 9.75                         | 9   |
| 30                        | 210                      | 19.5                         | 18  |

As an experiment progresses, POD5 files are produced for all reads. If you choose to basecall your data, these reads are used by the MinKNOW software to generate sequence data which is then stored in FASTQ files and/or BAM files.

# **Change log**

| Date                      | Version | Changes made  |
|---------------------------|---------|---|
| 20th<br>February<br>2024  | V17     | - In "Computer recommendations", the high performance option has been removed; references to adaptive sampling have been removed.       |
| 28th<br>September<br>2023 | V16     | <ul><li>A file size table has been added to "File types"</li><li>Minor corrections and clarifications throughout the document</li></ul> |

| Date                      | Version | Changes made  |
|---------------------------|---------|---|
| 23rd<br>August<br>2023    | V15     | <ul> <li>- A new section has been added: "Computer recommendations"</li> <li>- A new section has been added: "Connecting the MinION Mk1B to a computer without a USB Type-A port"</li> <li>- A new introductory paragraph has been added to "Host computer specification"</li> <li>- The disclaimer "The MinION device is CE marked using USB3.0. If a user wished to use USB-C, they may, but this invalidates the CE marking and may cause issues when using an adapter" has been removed.</li> </ul>   |
| 4th August<br>2022        | V14     | <ul> <li>In "Checklist", the specs for telemetry feedback, EPI2ME analysis, and software updates have been changed.</li> <li>"Networking explanations" has been removed.</li> <li>In "File types", information has been added about POD5 files and BAM files. The file size comparison table has been temporarily removed.</li> <li>The "Included software" section has been removed.</li> <li>Minor corrections and clarifications throughout the document.</li> </ul>   |
| 28th June<br>2023         | V13     | <ul> <li>Ubuntu 20.04 support has been included</li> <li>Host computer specifications have been updated from "recommended" to "required". The Intel CPU specification has been updated to requiring at least 4 cores/8 threads. Amperebased GPUs have been added to GPU recommendations.</li> <li>The requirement for solid-state storage has been reworded</li> <li>The amount of storage space required for 1 Gbyte of sequence data has been revised.</li> <li>In "Networking explanations", the AWS IP range has been updated to the global CloudFront IP range.</li> </ul> |
| 21st April<br>2022        | V12     | Updated the portal link from mirror.oxfordnanoportal.com to cdn.oxfordnanoportal.com in the host computer specifications section  |
| 10th<br>September<br>2021 | V11     | Removed erroneous compatibility with Ubuntu 20.04   |
| 8th<br>September<br>2021  | V10     | Added Windows 10 to the list of supported operating systems for GPU-enabled basecalling   |
| 22nd July<br>2021         | V9      | In the table of minimal requirements, two extra fields have been added: "Telemetry feedback, EPI2ME analysis", and "Software updates"   |
| 30th June<br>2021         | V8      | <ul> <li>Checklist at the top of the document has been removed</li> <li>Split IT requirements into Recommended (GPU) and Minimum (CPU)</li> <li>Updated operating systems recommendations for Windows (10), Mac OS (Mojave, Catalina) and Linux (18.04 and 20.04)</li> <li>Updated CPU recommendations to i7, i9, Xeon or better with 4+ cores</li> <li>Updated description and use of .fast5 files in the MinKNOW section</li> <li>Updated screenshot of the EPI2ME Agent</li> <li>Added table of .fast5 and FASTQ file sizes with different flow cell outputs</li> </ul>      |
| 23rd Dec<br>2020          | V7      | <ul> <li>Removed requirement for accessing IP address 106.187.40.102</li> <li>Updated the supported external NVIDIA GPUs to Tesla V100, Quadro GV100, Jetson TX2 and Jetson Xavier</li> </ul>   |
| 23rd Oct<br>2020          | V6      | Updated OSX compatibility   |
| 3rd July<br>2020          | V5      | Updated Fig 1 to 4.0.8 UI   |

| Date             | Version | Changes made   |
|------------------|---------|--|
| 18th May<br>2020 | V4      | Removed Bluetooth compatibility for MinIT              |
| Feb 2020         | V3      | More information about basecalling on GPUs using Guppy |