



Quick Reference

TDMx Amikacin

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Application

TDMx Amikacin

Developer

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Citation

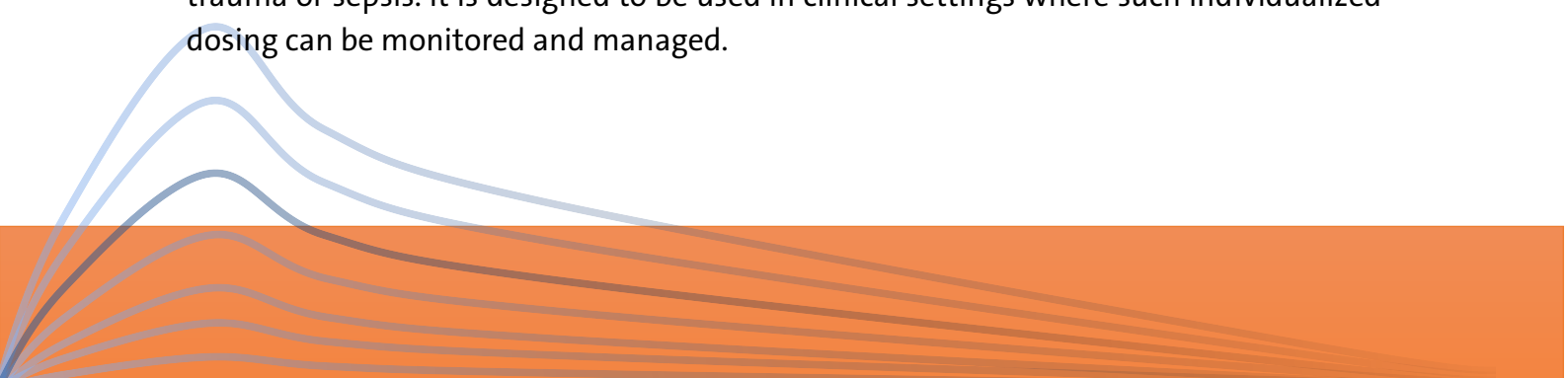
S.G. Wicha, M.G. Kees, A. Solms, I.K. Minichmayr, A. Kratzer, C. Kloft. TDMx – a novel web-based open-access support tool for optimising antimicrobial dosing regimens in clinical routine. *Int. J. Antimicrob. Agents*. 45: 442-444 (2015).

Purpose

The TDMx Amikacin software is intended for healthcare professionals to calculate individualized dosing regimens for the aminoglycoside antibiotic amikacin in critically-ill adult patients. The software aims to address the problem of high inter-patient variability in amikacin exposure achieved with standard dosing, which can result in concentrations that are not within the range needed to balance efficacy against nephro-/ototoxicity risk.

TDMx Amikacin utilizes patient-specific data such as demographics, trauma and sepsis status, laboratory values (serum creatinine), and, if available, previous dosing records and measured amikacin plasma concentrations, to determine individual pharmacokinetic parameters using the implemented population pharmacokinetic model (see [Select a model](#)). This computational task is designed to assist healthcare professionals in achieving a peak (C_{max}/MIC) >10 and $AUC_{24h}/MIC >70$ $mg \cdot h/L$ for efficacy, while keeping the trough (C_{min}) below the safety threshold appropriate to the dosing strategy used (<5 mg/L for extended-interval, <10 mg/L for traditional multiple-daily dosing — amikacin's own toxicity thresholds are higher than gentamicin's or tobramycin's).

TDMx Amikacin is intended for use in patients prescribed amikacin who require dose individualization for enhanced safety and efficacy, particularly critically-ill patients with trauma or sepsis. It is designed to be used in clinical settings where such individualized dosing can be monitored and managed.

A decorative graphic at the bottom of the page consisting of several overlapping, wavy lines in shades of blue and orange, creating a sense of movement and depth.

Intended users of the software are healthcare professionals (medical doctors, clinical pharmacists) who have expertise or training in therapeutic drug monitoring and pharmacokinetics. The operating environment for TDMx Amikacin should be healthcare facilities that have the capability to measure drug concentrations and monitor patient responses to antibiotic therapy.

TDMx Amikacin should not be used where it is not feasible to obtain the patient-specific data necessary for dose calculations, or where doing so would contravene the clinical judgement of the treating healthcare professional.

Disclaimer

TDMx is an educational tool and has been created for personal use only. The use of any result generated by TDMx is in any case the sole risk and responsibility of the TDMx user. The information provided by TDMx does not replace clinical judgement. Although TDMx has been validated carefully, there is no guarantee for the accuracy of the provided results. TDMx Amikacin has not been registered as a medical device; dosing decisions should not solely rely on TDMx and remain the responsibility of the treating physician.

Requirements

TDMx is fully tested on Google Chrome® and Mozilla Firefox® browsers. No installation is necessary — TDMx runs entirely in the browser.

Performance characteristics

TDMx was developed using reactive programming with a server-client architecture: TDMx reactively visualizes the user input such as dosing history, patient covariates or drug measurements. TDMx does not require any software to be installed locally.

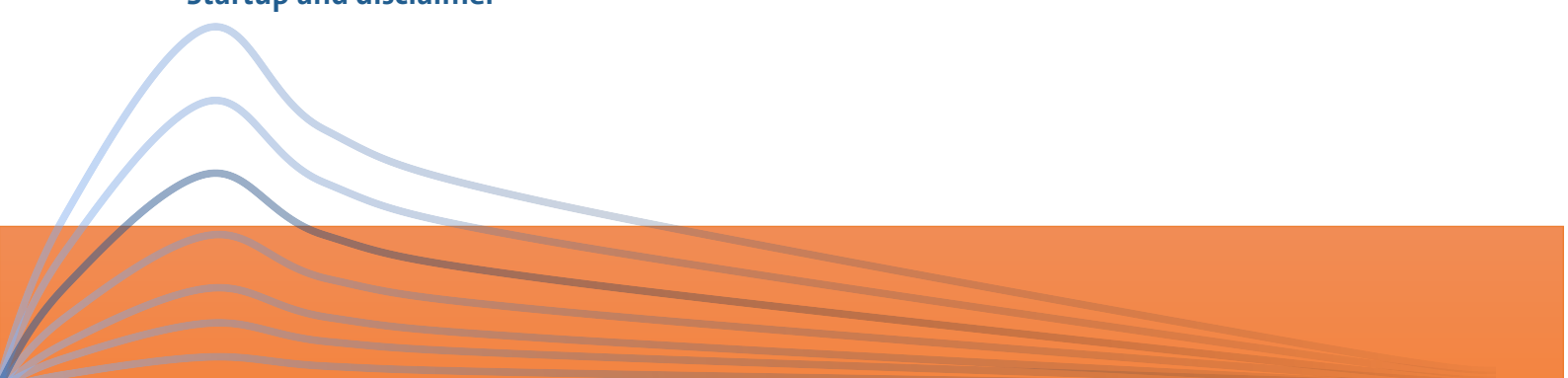
The population pharmacokinetic model implemented in TDMx Amikacin (see [Select a model](#)) was adapted from a published literature model (Romano et al. 1998) for critically-ill adults.

Launch TDMx from the web

Go to www.TDMx.eu

Navigate to the menu LAUNCH TDMX and select the module for amikacin.

Startup and disclaimer



During startup of the TDMx app, a welcome dialog is shown with the disclaimer above. Tick “I have read and accept the Disclaimer” and click “Start TDMx” to proceed (Figure 1). Loading a previously saved patient is done afterwards, directly in the main interface (see *Patient File*, below) — not as part of this startup step.

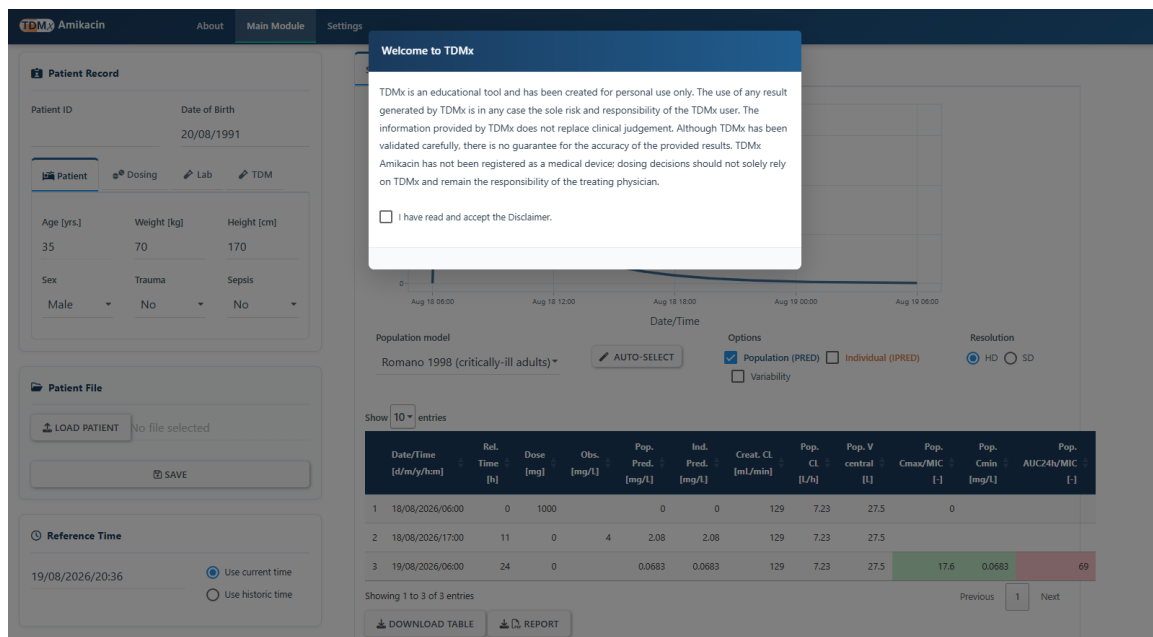


Figure 1: Startup dialog shown when launching TDMx Amikacin.

Main module

The main module consists of two panels: the patient panel (input, left) and the prediction/visualization panel (output, right) (Figure 2). Only one population pharmacokinetic model is implemented for amikacin (see [Select a model](#)), so there is no separate Model averaging tab here (unlike some other TDMx modules).

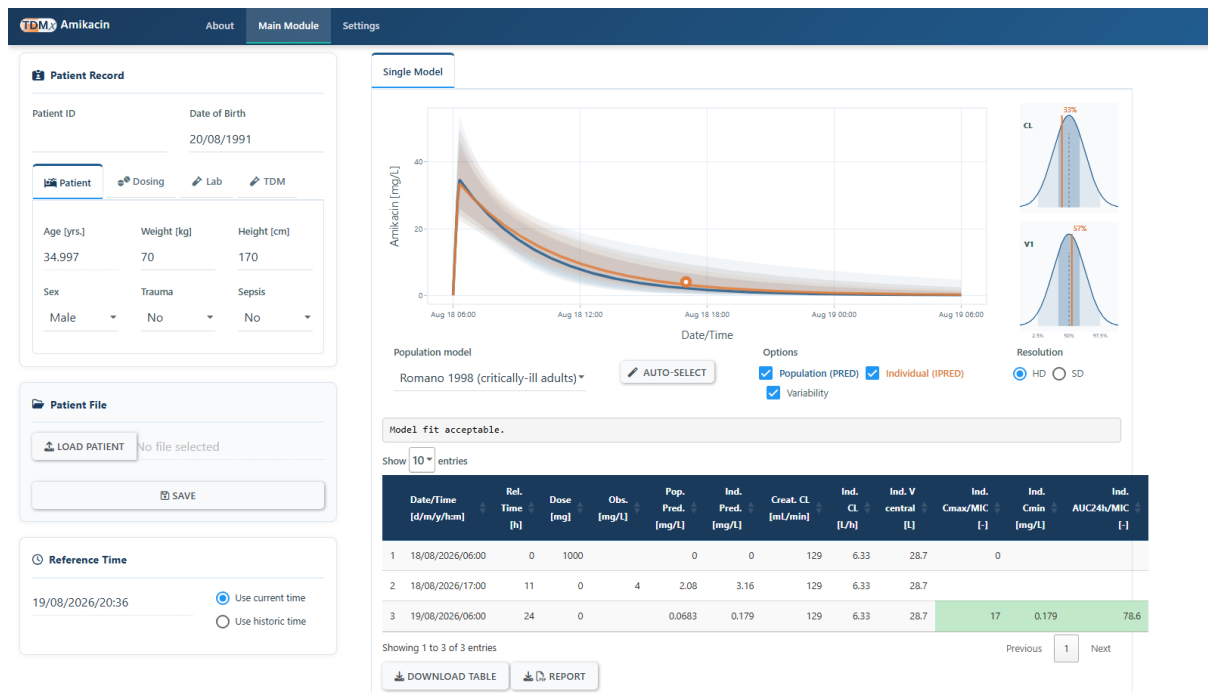


Figure 2: Main interface of the TDMx web-application, Patient tab, with Individual and Variability ticked.

Input panel (left column): contains three wells — *Patient Record* (patient ID and the Patient/Dosing/Lab/TDM tabs, described below), *Patient File* and *Reference Time*.

- *Patient File*: “LOAD PATIENT” opens a previously saved .tdmx file directly into the current session; “SAVE” downloads the current patient’s full record as a .tdmx file.
- *Reference Time*: the date/time TDMx treats as “now”. Switch to “Use historic time” to backdate a prediction, e.g. when reviewing a case retrospectively.

Tick any combination of the three Options next to the plot:

- Population (PRED, blue): Population-predicted concentration-time data using the dosing record and patient covariates. Observed concentration-time data are not used.
- Individual (IPRED, orange): Individually-predicted concentration-time data using the dosing record, patient covariates and the observed concentration-time data, via Bayesian estimation. Requires at least one TDM measurement.
- Variability: adds shaded prediction-interval bands (up to 95%) around the ticked Population and/or Individual line(s).

PK parameter density plot: whenever “Individual” is ticked and TDM data are available, a small panel of density curves (one per estimated PK parameter — CL and V1 only, since the Romano 1998 model has a single compartment) appears to the right of the main plot. Each curve is the population distribution of that parameter; the shaded band marks the 2.5th–97.5th percentile range and the vertical line marks where this patient’s individually estimated parameter falls, labelled with its percentile. A value near 50% indicates a typical patient; a value near the extremes (close to 2.5% or 97.5%) flags a potential outlier worth a closer look.

Model fit message: also shown once “Individual” is ticked, directly above the numerical table, is a one-line fit message:

- “Model fit acceptable.” (black) — both checks below passed.
- “Poor model fit.” (red) — the relative prediction error between observed and individually-predicted concentrations exceeds the configured threshold (default 40%).
- “...Patient parameters outside population distribution.” (appended, red) — one or more of this patient’s estimated PK parameters falls outside the model’s expected population range (outside its 95% interval by default).

The message turns red if either check fails, so it can also read “Model fit acceptable. Patient parameters outside population distribution.” — treat any red message as a prompt to inspect the fit and parameters carefully rather than accepting the prediction at face value.

Numerical output table:

- Date/Time: Time stamp of the event
- Dose: Administered dose [mg]
- Obs.: Observed concentration-time data [mg/L]
- Pop./Ind. Pred.: Population- or individually-predicted concentration-time data, respectively
- Creat. CL: Jelliffe creatinine clearance (body-surface-area corrected) — the actual covariate driving clearance in this model, not a display-only estimate
- Pop./Ind. CL/V central: Pharmacokinetic parameters of the population or individual model, respectively, considering patient covariates (and, for the

individual parameters, the observed concentration-time data). No V peripheral column — the Romano 1998 model has a single (central) compartment only.

- Pop./Ind. C_{max}/MIC, C_{min}, AUC_{24h}/MIC: Population- or individually-predicted peak-to-MIC ratio, trough concentration and 24-hour AUC-to-MIC ratio, shown for dosing rows only (MIC entered under the Lab tab)

Colour coding: the C_{max}/MIC, C_{min} and AUC_{24h}/MIC columns are each shaded green when within the target range configured under [Settings](#) and red otherwise — the same colours and thresholds used for the results table in the “Conventional Dosing (C_{max}-C_{min})” modal. This is a quick visual check, not a substitute for clinical judgement — always inspect the actual numbers.

Input panel

The input panel consists of four tabs: Patient, Dosing, Lab and TDM.

Patient

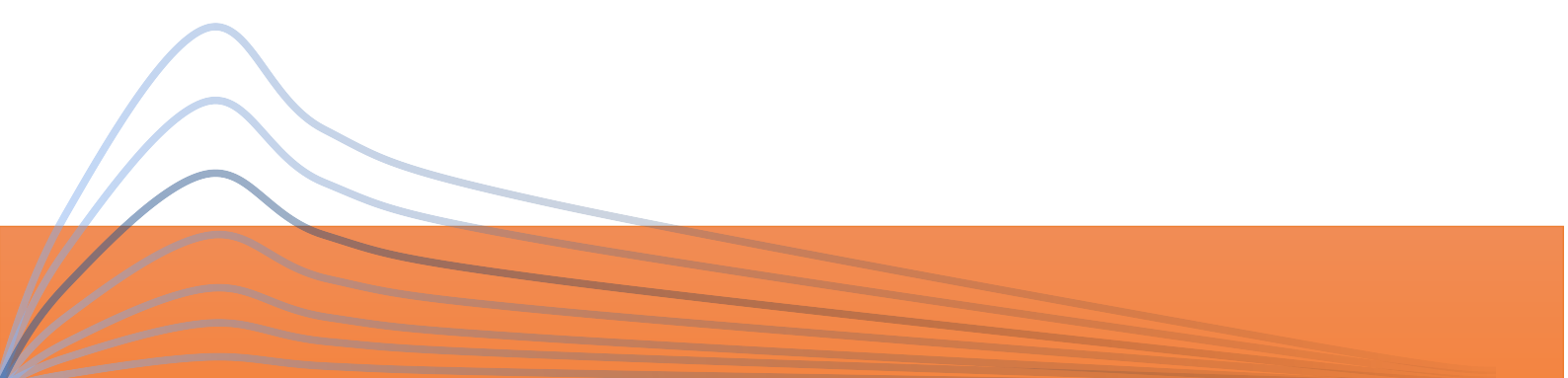
In this tab the user can specify the following (Figure 2):

- A name/identifier for this patient (Patient ID, above the tabs).
- Continuous covariates Age, Weight and Height, by either typing the value or using the up/down stepper.
- The categorical covariates Sex, Trauma and Sepsis, each selected from a dropdown menu.

Age, Weight and Height feed the (body-surface-area-corrected) Jelliffe creatinine clearance calculation that drives clearance. Trauma increases clearance; Sepsis increases the volume of distribution — both reflect physiological changes seen in critically-ill patients.

Dosing

Via the Dosing tab (Figure 3) the user can define all dosing events, and launch dose optimisation.



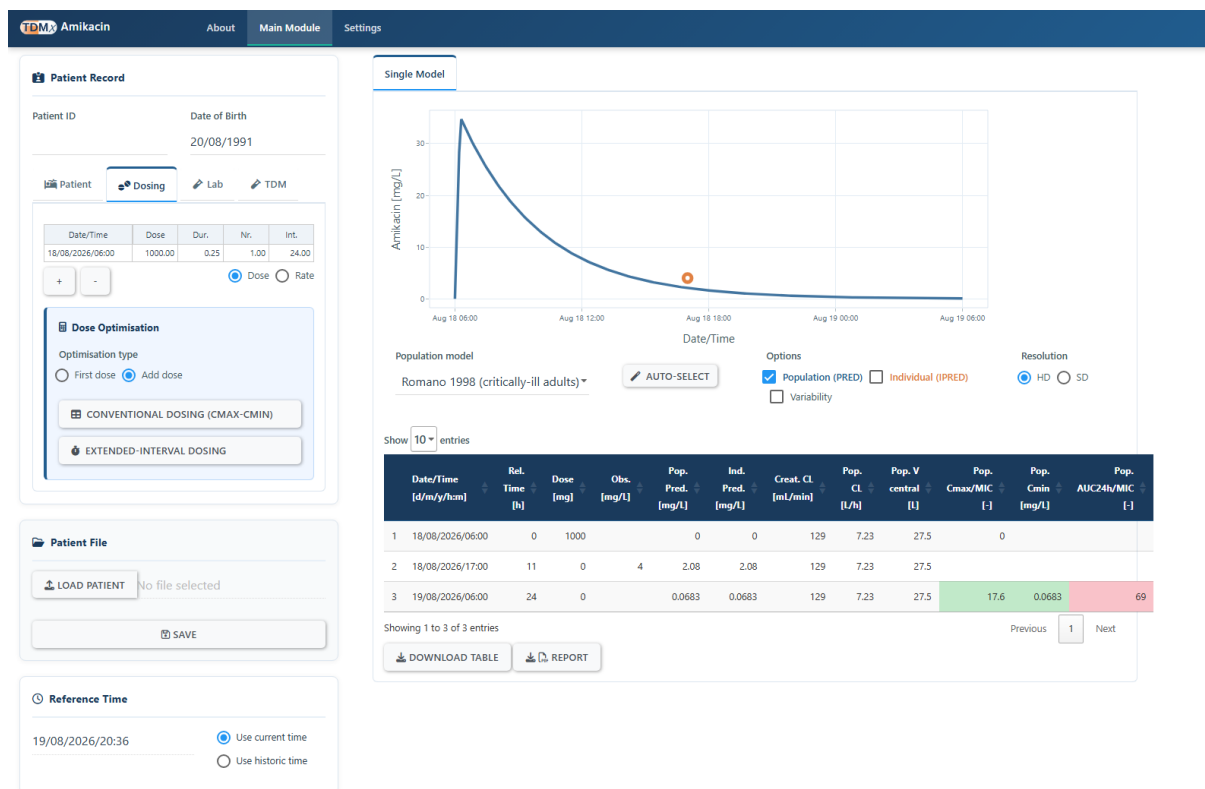


Figure 3: Input panel to edit dosing-related data.

A new dosing event can be added via “+”, while “-” removes the latest one. Specify the date (dd/mm/yyyy/hh:mm), the dose in mg (or, if switched to “Rate”, the infusion rate in mg/h) and the infusion duration (Dur.) in hours — amikacin is dosed intravenously into the central compartment only. If a dose is repeated, the number of repeats and the dosing interval (in hours) can be specified via Nr. and Int., respectively.

TDMx offers two dose-optimisation tools for amikacin; see [Dose optimisation](#) below.

Lab

In the Lab tab (Figure 4), the user can specify laboratory data obtained from the patient. These values are used directly as covariates in the population PK model and improve the prediction.

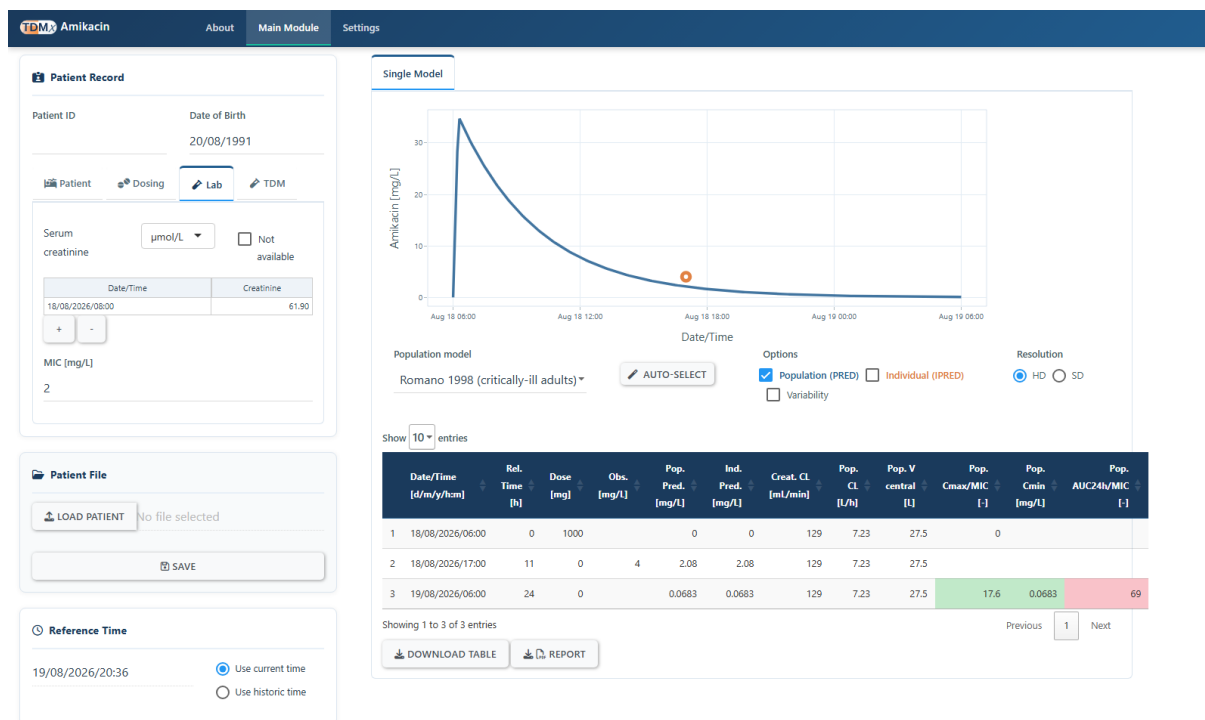


Figure 4: Input panel to enter laboratory data.

Available markers are Serum creatinine and the amikacin minimal inhibitory concentration (MIC). Specify the date of measurement as dd/mm/yyyy/hh:mm and the value in the stated (or an alternative) unit — TDMx automatically converts entered measurements between units. If no measurement is available, leave the field at its default value. For the MIC, provide an educated guess (default: 2 mg/L) if not (yet) available.

TDM

A new drug plasma concentration can be added via “+”, while “-” removes the latest measurement. Specify the date (dd/mm/yyyy/hh:mm) and the concentration in mg/L (Figure 5). Untick “Include” to exclude a sample from the Bayesian estimation without removing it from the plot.

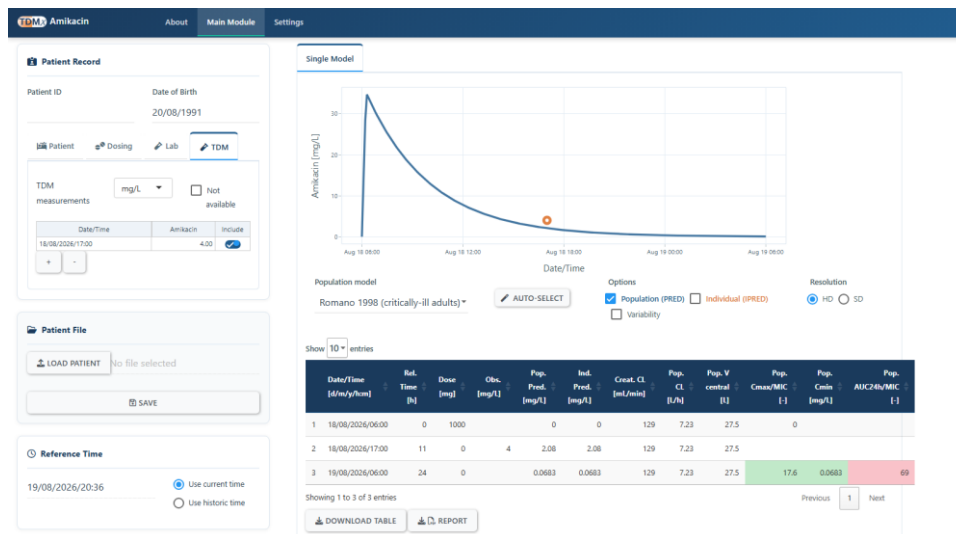


Figure 5: Input panel to enter TDM data.

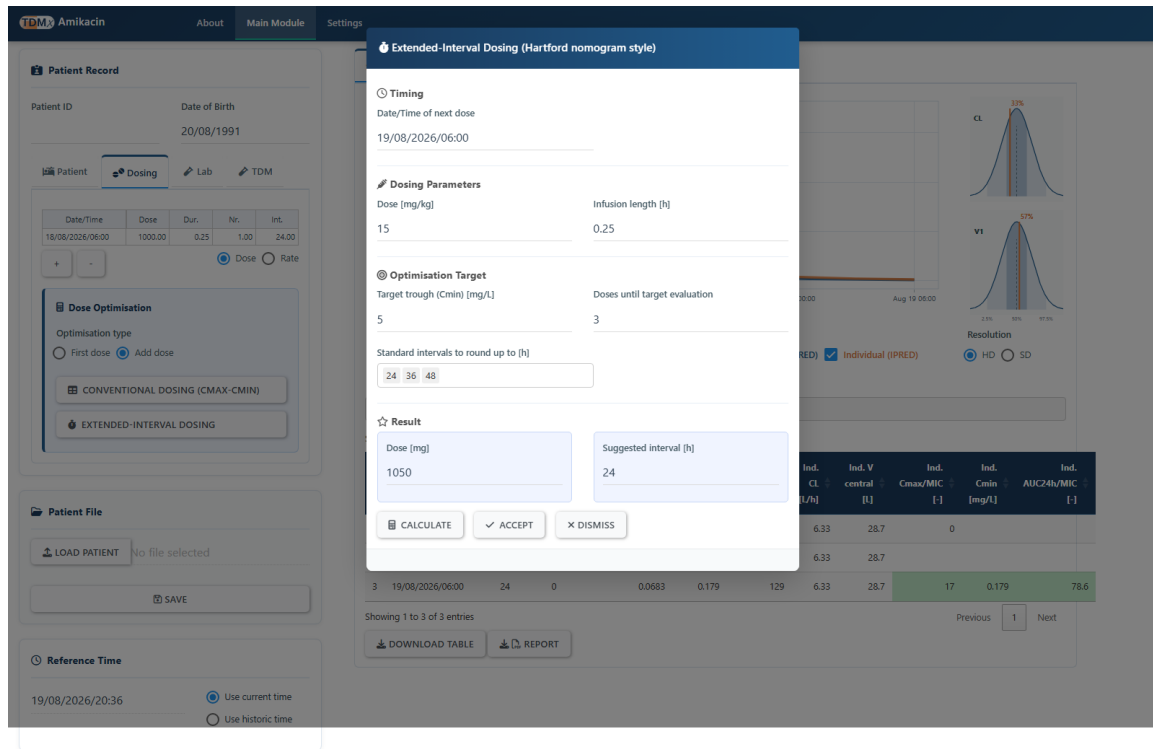
Dose optimisation

Both tools are opened from the Dosing tab and use the selected model, the patient's covariates and pathogen data (measured or assumed MIC), either with or without drug measurements. Optimising without drug measurements is useful to guide dosing before treatment starts or when no measurements are available; optimising with measurements supports ongoing therapy and is more precise, since individual PK parameters are used.

Note: if “First dose” is selected as the optimisation type, all previous dosing and TDM data will be removed. A warning is shown; accept it only if this is intended.

Extended-Interval Dosing

Clicking “EXTENDED-INTERVAL DOSING” opens the Hartford-nomogram-style dialog (Figure 6): a fixed, weight-based dose (default 15 mg/kg, editable — amikacin's typical extended-interval dose, higher than gentamicin/tobramycin's ~7 mg/kg) is given, and TDMx solves for the shortest dosing interval that keeps the predicted trough at or below the target (default 5 mg/L, editable) after the given number of repeated doses (*Doses until target evaluation*, default 3 — accounts for accumulation across repeated dosing rather than just after a single dose). The continuously-solved interval is rounded up to the shortest of the standard intervals selected that still reaches the target; if none do, the longest is used and a warning is shown. Click “CALCULATE” to solve, then “ACCEPT” to add the resulting dose and interval to the patient's dosing record — both the Dose and Suggested interval fields can be manually overridden before accepting.



The screenshot shows the TDMx Amikacin interface with the 'Extended-Interval Dosing (Hartford nomogram style)' dialog open. The background interface includes a 'Patient Record' section with fields for Patient ID, Date of Birth, and a 'Dosing' section with a table of dosing records. The dialog itself has several sections: 'Timing' with a date/time field, 'Dosing Parameters' with dose and infusion length fields, 'Optimisation Target' with target trough and doses until evaluation fields, and 'Result' with dose and suggested interval fields. There are 'CALCULATE', 'ACCEPT', and 'DISMISS' buttons at the bottom of the dialog. In the background, a nomogram graph is visible, showing two bell curves for CL and V1 with 50% and 97.5% markers. Below the graph is a table of results.

Ind. CL [l/h]	Ind. V central [l]	Ind. Cmax/MIC [l]	Ind. Cmin [mg/l]	Ind. AUC24h/MIC [l]
6.33	28.7	0		
6.33	28.7			
6.33	28.7	17	0.179	78.6

Figure 6: Extended-Interval Dosing dialog (Hartford nomogram style).

Conventional Dosing (Cmax-Cmin)

Clicking “CONVENTIONAL DOSING (CMAX-CMIN)” opens the dialog (Figure 7), which solves, for every one of a fixed set of candidate dosing intervals (8, 12, 18, 24, 36 and 48 h by default — always evaluated), the dose that reaches a target Cmax/MIC ratio (default 10, editable). The resulting Cmax/MIC, Cmin and AUC24h/MIC are read back and colour-coded against the Settings-tab thresholds. Click “CALCULATE” to solve every interval, select the most suitable row, and click “ACCEPT” to add it to the patient’s dosing record.

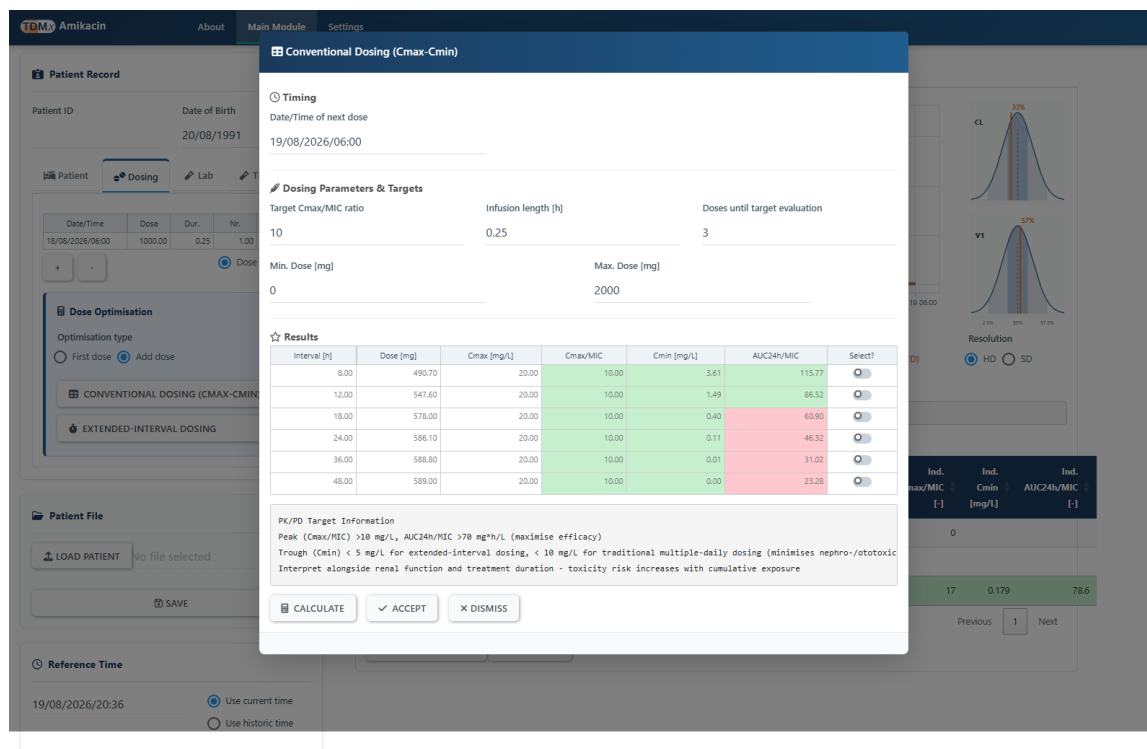


Figure 7: Conventional Dosing (Cmax-Cmin) dialog.

Both dialogs share a *Doses until target evaluation* field: the number of identical, repeated doses given (at the interval being solved, or at each candidate interval) before the trough/Cmax-Cmin/AUC24h-MIC is checked — accounts for accumulation across repeated dosing (higher = closer to steady state).

Select a model

Only one population pharmacokinetic model is currently implemented for amikacin:

Model	Population	Key covariates used
Romano 1998	Critically-ill adults	Weight, Height, Age, Sex, Serum creatinine, Trauma, Sepsis

The “AUTO-SELECT” control next to the Population model field has no effect while only one model is implemented — it exists for consistency with other TDMx modules that offer several candidate models.

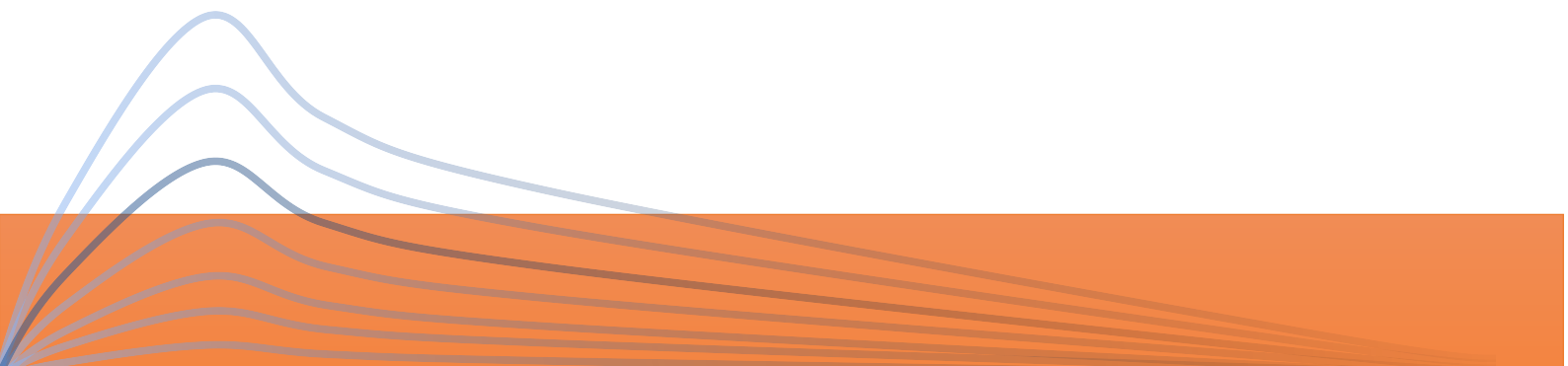
Model fit warnings

TDMx continuously checks the fit between the individual prediction and the entered TDM data against two thresholds, and reports the result as the fit message described under [Main module](#) (“Model fit acceptable.” / “Poor model fit.” / “...Patient parameters outside population distribution.”):

- **Residual check:** flags a poor fit if the relative prediction error, $|\text{predicted} - \text{observed}| / \text{mean}(\text{predicted}, \text{observed})$, exceeds 40% for any TDM sample (configurable).
- **Parameter check:** flags an outlier patient if any individually estimated PK parameter falls outside the 95% interval of that parameter’s population distribution (configurable), i.e. the same percentile shown by the PK parameter density plot.

If either check fails, treat the prediction with extra caution: inspect the plot and PK parameters carefully, and consider that a dosing regimen based on clinical judgement alone might be more appropriate for this patient than the one calculated by TDMx.

Settings



Amikacin

AboutMain ModuleSettings

SIMULATION RESOLUTION

HD Resolution [hrs]

SD Resolution [hrs]

0.1

2

MONTE CARLO SIMULATIONS

Prediction interval method:

☒ Delta-Method

☐ Monte Carlo

Replicates for visualisation

50

250

1,000

Replicates for dose prediction intervals

100

1,000

Variability components in MC simulation:

☒ Inter-individual

☒ Inter-occasion

☐ Residual

MODEL AVERAGING

Averaging/selection criterion:

Sum of squared residuals (SSE)

PK TABLE COLOUR CODING

Reference ranges used to colour-code target columns (green/red) in the PK table.

Target C_{max}/MIC [-]

Target trough (C_{min}) [mg/L]

0

10

50

0

5

15

Target AUC_{24h}/MIC [-]

0

10

200

SHARE

Shareable link with patient data encoded in URL:

http://127.0.0.1:9032/?data_json=K7BNk2dose_TableK2K3AK5BN7BNk2DateTImeK2K3AK21BNk2F0BNk2F20k2G

Figure 8: Settings tab.

To adjust the speed/performance and behaviour of TDMx:

- **Simulation Resolution:** separate time resolutions (in hours) for the high-definition (HD) and standard-definition (SD) plot/table output.
- **Monte Carlo Simulations:** choose the prediction-interval method — Delta-Method (fast, analytical approximation, default) or Monte Carlo (simulation-based) — and, for the latter, the number of replicates used for visualisation and for dose prediction intervals respectively. Variability components (Inter-individual, Inter-occasion, Residual) can be included or excluded from the simulation individually.

- **PK Table Colour Coding:** the target C_{max}/MIC ratio (default 10), target trough (C_{min}, default 5 mg/L) and target AUC_{24h}/MIC (default 70) used to colour-code the main PK table and the “Conventional Dosing (C_{max}-C_{min})” results table.
- **Share:** generates a shareable link with the current patient’s data encoded directly in the URL. This lets a colleague open the same patient record in their own browser session without needing to exchange a *.tdmx file.

Abbreviations

Abbreviation	Meaning
Ind.	Individual
HD	High definition
SD	Standard definition (lower time resolution)
PK	Pharmacokinetics
Pred.	Prediction(s)
Obs.	Observed plasma concentration
TDM	Therapeutic drug monitoring
CL	Clearance
CRCL	Creatinine clearance
C _{max}	Peak concentration
C _{min}	Trough concentration
AUC	Area under the curve
MIC	Minimum inhibitory concentration