## Molecular computer experiment

Also pseudoexperiment

| REAL EXPERIMENT | COMPUTER EXPERIMENT |
| :--- | :--- |
| Record everything in a lab notebook | Record everything in a lab notebook |
| Choose method (device, assay) | Choose method (MD, MC,...) |
| Build the experimental apparatus <br> (from parts) | Download/buy/write a computer program (blocks of <br> code) |
| Purchase chemicals, synthetise if not <br> available | Get a force field, fit/calculate parameters if not avail- <br> able (e.g., partial charges) |
| Prepare the experiment | Prepare initial configurations, etc. |
| Perform the experiment, watch <br> what's going on | Run the code, observe time development, control <br> quantities, etc. |
| Analyse and calculate | Calculate mean values (with error estimates) |
| Clean the laboratory | Make backups, erase temporary files |

MD or MC?
Often, MC and MD can be applied to similar systems.
MD
realistic models, complex molecules (bonds, angles...)
condensed matter in general (fluids, solutions; biochemistry)
kinetic quantities (diffusivity, viscosity....)
better parallelization, more packages available
MC
simple qualitative models (lattice, hard-sphere-like)
dilute systems
critical phenomena
fluid equilibria
overcoming barriers, exchange of molecules, etc. is easier with MC
less efficient parallelization, fewer packages available
Is it correct? $\quad 3 / 15$

## Systematic errors:

- inaccurate molecular model (force field)
- neglected quantum effects, neglected many-body forces ...
- small sample (finite-size effects)
insufficient time scale (long correlations, bottleneck problems)
. method problems: integration errors (too long timestep), inappropriate thermostat/barostat, not equilibrated enough, inaccurate treatment of Coulomb forces. .
Random (stochastic, statistical) errors* are essential in stochastic methods
time-correlated
- can be decreased by long calculations

Uncertainty (in metrology) includes critical assessment of both the systematic and random errors*

* different terminology in different fields (mathematical statistics, metrology, physics, chemistry)


## Simulation methodology

[sleep 3;simul/spceE.sh] 4/15

## Start (initial configuration):

- experimental structure (biomolecules)
crystal $\rightarrow$ liquid (melt), gas $\rightarrow$ liquid (shrink); Packmol
- random configuration (overlaps of molecules = problem in MD) problem for "ill-defined" models (TIP4P etc.)
- lattice models: crystal/chaos
- MD: velocities $=$ Maxwell-Boltzmann (approximation enough)
- Equilibration $\rightarrow$ watch graphically (convergence/time profile)
- Measuring the quantities of interest incl. estimates of errors



## Calculations

[../simul/ar/showdrop.sh] $5 / 15$
$506 / 3$
Example. We simulate an argon droplet in a periodic cubic simulation cell. Let us have $N=1000$ atoms and temperature $T=85 \mathrm{~K}$. The distance between surfaces of periodic images of droplets should be equal to the droplet diameter. Calculate the size of the box in $\AA \AA$. Argon density is $\rho=$ $1.4 \mathrm{~g} \mathrm{~cm}^{-3}$, molar mass $M(\mathrm{Ar})=40 \mathrm{~g} / \mathrm{mol}$.
molar volume: $V_{\mathrm{m}}=M / \rho$
volume per 1 atom: $V_{1}=V_{\mathrm{m}} / N_{\mathrm{A}}$
volume of $N$ atoms: $V=N V_{1}=N M / \rho N_{A}$
$=1000 \cdot 0.040 \mathrm{~kg} \mathrm{~mol}^{-1} /\left(1400 \mathrm{~kg} \mathrm{~m}^{-3} \cdot 6.022 \times 10^{23} \mathrm{~mol}^{-1}\right)$
$=4.744 \times 10^{-27} \mathrm{~m}^{3}$
droplet radius: $\frac{4}{3} \pi R^{3}=V \Rightarrow R=2.24 \times 10^{-9} \mathrm{~m}$
box size: $\underline{L=90 \AA}$


## One more example

Example. Consider a globular protein of molecular weight of 20 kDa . The density of the protein is $1.35 \mathrm{~g} \mathrm{~cm}^{-3}$. Calculate the approximate protein diameter.

$$
m=\frac{20 \mathrm{~kg} \mathrm{~mol}^{-1}}{6.022 \times 10^{23} \mathrm{~mol}^{-1}}=3.32 \times 10^{-23} \mathrm{~kg}
$$


or $1 \mathrm{Da}=1 \mathrm{~g} \mathrm{~mol}^{-1} / \mathrm{N}_{\mathrm{A}}=1.6605 \times 10^{-27} \mathrm{~kg}$ (atomic mass unit)
$m=20000 \times 1.6605 \times 10^{-27} \mathrm{~kg}=3.32 \times 10^{-23} \mathrm{~kg}$

$$
V=\frac{m}{\rho}=\frac{3.32 \times 10^{-23} \mathrm{~kg}}{1350 \mathrm{~kg} \mathrm{~m}^{-3}}=2.46 \times 10^{-26} \mathrm{~m}^{3}
$$

$$
\frac{4 \pi}{3} r^{3}=\frac{\pi}{6} d^{3}=V
$$

$$
d=\sqrt[3]{\frac{6 V}{\pi}}=\sqrt[3]{\frac{6 \cdot 2.46 \times 10^{-26} \mathrm{~m}^{3}}{\pi}}=3.61 \times 10^{-9} \mathrm{~m} \doteq \underline{3.6 \mathrm{~nm}}=36 \AA
$$

## Measurements

Trajectory $=$ sequence of configurations (MD: in time)

## Convergence profile:

time development of a quantity (time profile, -) problems better seen

- cumulative (running average, - )
can estimate the inaccuracy


## Type of statistical treatment:

averaged values ( $\leftarrow$ ergodic hypothesis)


- less often fluctuations


## Type of quantity:

mechanical (temperature, pressure, internal energy, order parameters...)

- entropic ( $S, F, \mu, \ldots$ )
- structure (correlation functions, number of neighbors, analysis of clusters...)
auxiliary or control quantities (order parameters, integrals of motion in MD)


## Random errors

quantity $=$ (estimate of the mean value) $\pm$ (estimate of the error)
Arithmetic average (example of a statistic, also statistical functional, estimator, in metrology measurement function):

$$
\bar{X}=\frac{1}{m} \sum_{i=1}^{m} x_{i} \quad \begin{aligned}
& \text { statistic }=\text { estimator } \\
& \text { statistics }=\text { field of mathematics }
\end{aligned}
$$

Standard error $=$ standard deviation of the statistic, usually denoted as $\sigma$

$$
\sigma_{x}=\sqrt{\left\langle(\bar{x}-\langle x\rangle)^{2}\right\rangle}
$$

For uncorrelated (independent) $X_{i}$ and large $m, \bar{X}$ has Gaussian normal distribution
The estimate of the standard error of the arithmetic average of uncorrelated data:

$$
\sigma_{X}^{\text {estim }}=\sqrt{\frac{\sum_{i=1}^{m} \Delta x_{i}^{2}}{m(m-1)^{\prime}}} \quad \text { where } \Delta x_{i}=x_{i}-\bar{X}
$$

## Customs and bad habits

How the uncertainty of measured quantities are expressed in different fields:
Physics: $Q=123.4 \pm 0.5 \equiv 123.4(5) \equiv 123.45 \quad \AA$
$0.5=\sigma(Q)=$ (estimated) standard error/uncertainty of statistic $Q$ (e.g., $Q=\bar{X}$ ), also: standard deviation (meaning of the average or other statistic)
loosely: (estimated) error/uncertainty, standard deviation, error margin, error bar,
In case normal distribution, it holds $\langle Q\rangle \in 123.4 \pm 0.5$ with probability $68 \%$

- Biology, economy, politology, engineering, pharmacology: $Q=123.4 \pm 1.0 ~ \mathrm{~K} \pm 1.0=$ $\pm 2 \sigma(Q)=$ confidence interval at (confidence) level $95 \%$
looselyn: $\pm 1.0=$ confidence interval, $1.0=$ error/uncertainty,
In case normal distribution, it holds $\langle Q\rangle \in 123.4 \pm 1.0$ with probability $95 \%$
- Chemistry: often ignored; if given, nobody knows the confidence level
, „Physical certainty" starts at $\pm 5 \sigma_{X}$ (confidence level 0.999999 43)
The type of error/uncertainty must be always specified
$\alpha=$ significance level, often $5 \%$
$1-\alpha=$ confidence level, often $95 \%$


## Analysis of time series and error estimation

## Problem: correlations

block method: $\bar{X}_{j}=\frac{1}{B} \sum_{i=1}^{B} X_{i+(j-1) B}$

- analysis of correlations $\Rightarrow$

$$
\sigma_{X}=\sqrt{\frac{\sum_{i=1}^{m} \Delta X_{i}^{2}}{m(m-1)}(1+2 \tau)} \quad \tau=\sum_{k=1}^{\infty} c_{k} \quad c_{k}=\frac{\left\langle\Delta X_{0} \Delta X_{k}\right\rangle}{\left\langle(\Delta X)^{2}\right\rangle}
$$

$\mathrm{MC}: c_{k}$ is monotonously decreasing [ex.: $c_{k}=\sum_{\lambda \neq 1} c_{\lambda} \lambda^{k}, \lambda \in(-1,1)$ ]
MD: $c_{k} \rightarrow c(t)$ (time autocorrelation function): damped oscillations

- even better = both approaches combined:
first to block a bit, then $\tau \approx c_{1}$
- from running average (roughly $\approx 10$ blocks):

$$
\sigma_{X}^{\text {estim }} \approx 0.6\left[\max _{2 \text { nd half }}(X)-\min _{2 \text { nd half }}(X)\right]
$$

or to be on the safe side (this formula is approximate):
$\operatorname{err} x \approx \max _{2 \text { nd half }}(x)-\min _{2 \text { nd }}$ half $(X)$
$\Rightarrow\langle X\rangle \in\left(\bar{X}-\mathrm{err}_{X}, \bar{X}+\right.$ err $X$ ) with probability $\approx 85 \%$ (for long enough time series)

where the variance, or fluctuation, is defined by $\operatorname{Var} X=\left\langle(X-\bar{X})^{2}\right\rangle$


Error analysis - division and multiplication

Example. Calculate 3.46(7)/0.934(13).
fraction: 3.46/0.934 $=3.704$
rel. error $=\sqrt{\left(\frac{0.07}{3.46}\right)^{2}+\left(\frac{0.013}{0.934}\right)^{2}}=0.0246$
abs. error $=3.704 \times 0.0246=0.091$
$3.46(7) / 0.934(13)=3.70(9)$ (or rounded up: 3.70(10))

## Error analysis

Error of function $f$ of a variable with error is (linearized; i.e., for small $\sigma$ ):

$$
f\left(x \pm \sigma_{x}\right)=f(x) \pm f^{\prime}(x) \sigma_{x}
$$

$$
\ln \left(x \pm \sigma_{x}\right)=\ln x \pm \frac{\sigma_{x}}{x}, \quad \exp \left(x \pm \sigma_{x}\right)=\exp x \pm \sigma_{x} \exp x, \quad \frac{1}{x \pm \sigma_{x}}=\frac{1}{x} \pm \frac{\sigma_{x}}{|x|^{2}}
$$

Example. Calculate the activity of $\mathrm{H}^{+}$from $\mathrm{pH}=2.125(5)$. activity:

$$
a_{\mathrm{H}^{+}}=10^{-2.125}=\exp (-2.125 \times \ln 10)=0.00750
$$

error Method 1:

$$
\sigma=0.005 \times \ln 10 \times a=0.000086
$$

error Method 2:

$$
\sigma=\left|10^{-2.125}-10^{-2.125-0.005}\right|=0.000087
$$

activity with error (uncertainty) estimate:
$a_{\mathrm{H}^{+}}=0.00750(9)$

