

Automatic differentiation for error analysis

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References:

- U. Wolff, "Monte Carlo errors with less errors". Comput.Phys.Commun. 156 (2004) 143-153.
- F. Virotta, "Critical slowing down and error analysis of lattice QCD simulations." PhD thesis.
- Stefan Schaefer, Rainer Sommer, Francesco Virotta, "Critical slowing down and error analysis in lattice QCD simulations". Nucl.Phys.B 845 (2011) 93-119.
- A. Ramos, "Automatic differentiation for error analysis of Monte Carlo data". Comput.Phys.Commun. 238 (2019) 19-35.
- M. Bruno, R. Sommer, In preparation.

Software: <https://igit.ific.uv.es/alramos/aderrors.jl>



DATA ANALYSIS IN LATTICE QCD

From simulations to the proton mass

- ▶ Discretize space-time in a lattice of spacing a
- ▶ Use Monte Carlo techniques to generate “configurations” (representatives of the QCD vacuum)
- ▶ “Measure” correlation functions in these configurations

$$aM_p, af_\pi, af_k, af_+^{B \rightarrow D}(q^2), aM_\pi, \dots$$

- ▶ Detailed analysis to extrapolate results to the physical world:

$$a \rightarrow 0, L \rightarrow \infty, m_q \rightarrow m_q^{\text{phys}}, \dots$$

- ▶ Our result (i.e. $M_p = 943(17)\text{MeV}$, $V_{cb} = 0.039(3)$, ...) is a (complicated) **function** of the “measured” quantities in **several** ensembles (values of a, L, m_q)

$$M_p = F(aM_p, af_\pi, af_k, aM_\pi, \dots)$$

- ▶ This talk: techniques to determine uncertainty in **derived observables**

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LINEAR ERROR PROPAGATION

Seems trivial...

$$Z = F(X) \implies \delta Z = \frac{dF}{dX} \delta X$$

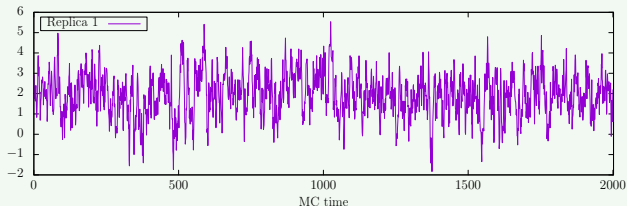
$$X = 1.0 \pm 0.1 \implies Z = \sin(X) = 0.841 \pm 0.054$$

But we need to be careful with

- ▶ Correlations between data
- ▶ What happens when F is an iterative algorithm

F = Fit data (x, y) to model f

- ▶ Final error... Where does it come from?
- ▶ What to do when data is not $X = 1.0 \pm 0.1$, but



WELCOME TO ADerrors.jl

- ▶ Exact linear error propagation, even in iterative algorithms. Thanks to Automatic differentiation (ForwardDiff.jl)
- ▶ Handles data from any number of ensembles (i.e. simulations with different parameters/data from different sources).

Code (Julia)

```
1 julia> import Pkg
2 (v1.X) pkg> add https://igit.ific.uv.es/alramos/bdio.jl
3 (v1.X) pkg> add https://igit.ific.uv.es/alramos/aderrors.jl
```

A CALCULATOR WITH UNCERTAINTIES

```
Code (Julia)
1 julia> x = uwreal([12.31, 0.23], "Experiment A") # x = 12.31(23) from some experiment
2 12.31 (Error not available... maybe run uwerr)
3
4 julia> y = uwreal([4.22, 0.12], "Experiment B") # x = 4.22(12) from some other experiment
5 4.22 (Error not available... maybe run uwerr)
6
7 julia> z = x + y
8 16.53 (Error not available... maybe run uwerr)
9 julia> uwerr(z)
10
11 julia> println(z) # sqrt(0.23^2 + 0.12^2) = 0.25942243542145693
12 16.53 +/- 0.25942243542145693
13
14 julia> details(z)
15 16.53 +/- 0.25942243542145693
16 ## Number of error sources: 2
17 ## Number of MC ids      : 0
18 ## Contribution to error :
19 #                          Ensemble [%]      [MC length]
20 #                          Experiment A  78.60      -
21 #                          Experiment B  21.40      -
22
23 julia> zero_error = sin(z) - ( sin(x)*cos(y) + cos(x)*sin(y) )
24 -6.661338147750939e-16 (Error not available... maybe run uwerr)
25 julia> uwerr(zero_error)
26
27 julia> println(zero_error) # ADerrors keeps correlations!
28 -6.661338147750939e-16 +/- 1.7281005654554058e-16
```

ADerrors.jl ALLOWS TO INPUT CORRELATED DATA

```
Code (Julia)
1 julia> avg = [16.26, 0.12, -0.0038];
2 julia> Mcov = [0.478071 -0.176116 0.0135305
3               -0.176116 0.0696489 -0.00554431
4               0.0135305 -0.00554431 0.000454180];
5
6 julia> p = cobs(avg, Mcov, "Correlated data")
7 3-element Array{uwreal,1}:
8 16.26 (Error not available... maybe run uwerr)
9 0.12 (Error not available... maybe run uwerr)
10 -0.0038 (Error not available... maybe run uwerr)
11
12 julia> uwerr.(p);
13 julia> p
14 3-element Array{uwreal,1}:
15 16.26 +/- 0.6914267857119798
16 0.12 +/- 0.2639107803785211
17 -0.0038 +/- 0.021311499243366245
18 julia> cov(p)
19 3x3 Array{Float64,2}:
20 0.478071 -0.176116 0.0135305
21 -0.176116 0.0696489 -0.00554431
22 0.0135305 -0.00554431 0.00045418
23
24 julia> z = p[1] + p[2] + sin(p[3]); # Correlations are propagated
25 julia> uwerr(z)
26 julia> z
27 16.37620000914533 +/- 0.4603415450741195
```

WORKS WITH ITERATIVE ALGORITHMS: ROOT OF $f(x) = a \cos(b \sin(x)) - x$

```
Code (Julia)
1 julia> a = uwreal([1.34, 0.12], "Data 01") # a = 1.34 +/- 0.12
2 julia> b = uwreal([1.34, 0.12], "Data 02") # b = 1.34 +/- 0.12
3
4 julia> x0 = uwreal([0.5,0.5], "Initial position") # x0 = 0.5 +/- 0.5
5 julia> while true # This is just newton method
6     val = a*cos(b*sin(x0)) - x0
7     der = -a*b*sin(b*sin(x0))*cos(x0) - 1.0
8     x1 = x0 - val/der
9     if (abs(value(x0) - value(x1))<1.0E-10)
10        break
11    else
12        x0 = x1
13    end
14 end
15
16 julia> uwerr(x1)
17
18 julia> details(x1)
19 0.7838003744331717 +/- 0.056992589665847124
20 ## Number of error sources: 3
21 ## Number of MC ids      : 0
22 ## Contribution to error :           Ensemble [%]      [MC length]
23 #                         Data 02 63.25             -
24 #                         Data 01 36.75             -
25 #                         Initial position 0.00             -
```


AWESOME... BUT I DO NOT WANT TO RE-CODE LEVENBERG-MARQUARDT

AD allows to propagate errors to fit parameters

- ▶ Find p_i ($i = 1, \dots, N_{\text{param}}$) that minimize

$$\chi^2(p_i; d_a), \quad p_i (i = 1, \dots, N_{\text{param}}), \quad d_a (a = 1, \dots, N_{\text{data}}).$$

with d_a some MC data.

- ▶ χ^2 is minimum at \bar{p}_i for the central values of the data \bar{d}_a
- ▶ How much changes the values of the parameters ($\bar{p}_i \rightarrow \bar{p}_i + \delta p_i$) that minimize χ^2 when the data is shifted $\bar{d}_a \rightarrow \bar{d}_a + \delta d_a$?

$$\frac{\delta p_i}{\delta d_a} = - \sum_{j=1}^{N_{\text{param}}} (H^{-1})_{ij} \partial_j \partial_a \chi^2 \Big|_{(\bar{p}_i; \bar{d}_a)}.$$

with

$$H_{ij} = \partial_j \partial_i \chi^2 \Big|_{(\bar{p}_i; \bar{d}_a)},$$

- ▶ Error propagation only requires derivatives of χ^2 at the minima (we do not care how you arrived there!)

EXACT ERROR PROPAGATION IN FITS

```
Code (Julia)
1  ...
2  julia> lm, csq = FFfit_funcs(fp, f0, 4, 4, nlatt, FLAGp); # Definition of fit functions
3  julia> uwdt
4  9.5    1.05004 +/- 0.039482274503883384
5  11.6   1.17597 +/- 0.04580916938779833
6  8.5    1.00648 +/- 0.012263563919187603
7  ...
8  julia> r = optimize(xx -> lm(xx, value.(uwdt)), zeros(9), LevenbergMarquardt())
9  * Algorithm: LevenbergMarquardt
10 * Minimizer: [0.03562259277485376, -0.2733271920349287, 1.6897944291757172, -89.38349668726151, 0.0
11 * Sum of squares at Minimum: 4.901151
12
13 julia> fitp, cexp = fit_error(csq, r.minimizer, uwdt) # Error propagation with ADerrors
14 julia> uwerr.(fitp) # The fit parameters are normal uwreal variables
15 julia> print("Vcb: ") # The last fit parameter is Vcb
16 julia> details(fitp[end])
17 Vcb: 0.04032176066923957 +/- 0.0021953643904483967
18 ## Number of error sources: 24
19 ## Number of MC ids      : 0
20 ## Contribution to error :
21 #           Ensemble [%]      [MC length]
22 #           BaBar 00000001  31.84      -
23 #           BaBar 00000002  23.33      -
24 #           BaBar 10b 00000001  16.30      -
25 #           BaBar 10b 00000002  11.80      -
26 #           BaBar 00000003  10.13      -
27 #           Lattice form factors 00000003  2.54      -
28 ...
```

HANDLING MONTE CARLO DATA

- Generate a random walk in the interval $[-1, 1]$

Code (Julia)

```
1 julia> # Generate some correlated data
2     eta = randn(10000);
3 julia> x = Vector{Float64}(undef, 10000);
4 julia> x[1] = 0.0;
5 julia> for i in 2:10000 # This is just a random walk in [-1,1]
6     x[i] = x[i-1] + 0.2*eta[i]
7     if abs(x[i]) > 1.0
8         x[i] = x[i-1]
9     end
10 end
```

- Input a Monte Carlo history as uwreal. Correlations handled automatically

Code (Julia)

```
1 julia> xp2 = uwreal(x.^2, "Random walk ensemble in [-1,1]")
2 0.3533602504472119 (Error not available... maybe run uwerr)
3
4 julia> xp4 = uwreal(x.^4, "Random walk ensemble in [-1,1]")
5 0.2197572322981959 (Error not available... maybe run uwerr)
6
7 julia> cov([xp2, xp4])
8 2×2 Array{Float64,2}:
9  8.12723e-5  6.57753e-5
10 6.57753e-5  5.28194e-5
```

CONCLUSIONS

Error propagation with automatic differentiation

- ▶ **Robust:** If central values are correctly computed, errors will be correctly propagated.
- ▶ **Faster** and more **accurate** than alternatives based on resampling (i.e. jackknife, bootstrap)
- ▶ Correlations taken care of automatically
- ▶ **Exact** Error propagation in iterative algorithms: errors in fit parameters
- ▶ Bookkeeping of the contribution of each source to the error of a variable
- ▶ Autocorrelations of Monte Carlo data handled robustly

If you want to understand the theoretical ideas

A. Ramos, "Automatic differentiation for error analysis of Monte Carlo data". Comput.Phys.Commun. 238 (2019) 19-35.

Free implementation available

<https://igit.ific.uv.es/alramos/aderrors.jl>

(also fortran implementation available if interested)