Introduction to Automatic Differentiation

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- Automatic Differentiation (AD)
 - Definition by example
 - Forward and reverse mode
 - Scalar and vector mode
- AD implementation
 - Source transformation and operator overloading
 - Reverse mode example
 - Tools
- Alternatives to AD
 - Divided differences, Complex-Variable method
 - Symbolic and manual differentiation
- Summary





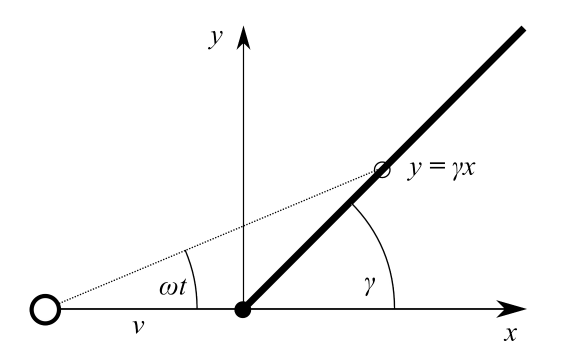
• Automatic or Algorithmic Differentiation (AD)

- Given a numeric program, that implements function ${\cal F}$
- AD creates a new program that computes F', the first order derivative of F
- And sometimes also the higher order derivatives F'', F''', $F^{\rm IV}$, etc.





• Consider the beam of a lighthouse rotating with angular velocity ω as it runs along a quay with slope γ at distance v, as a function of time t





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• The coordinates of the point where the light hits the quay are given by

$$x = \frac{\nu \tan(\omega t)}{\gamma - \tan(\omega t)} \qquad \qquad y = \frac{\gamma \nu \tan(\omega t)}{\gamma - \tan(\omega t)}$$

• A program implementing this function

$$v_1 = \omega * t;$$

$$v_2 = \tan(v_1);$$

$$v_3 = \gamma - v_2;$$

$$v_4 = \nu * v_2;$$

$$x = v_4/v_3;$$

$$y = \gamma * x;$$

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- Program code can be mechanically differentiated
 - Differentiate each statement and insert it before the original statement $\delta v_1 = \delta \omega * t + \omega * \delta t$:

 $v_1 = \omega * t;$ $v_1 = \omega * t$: $\delta v_2 = \delta v_1 / \cos^2(v_1);$ $v_2 = \tan(v_1);$ $v_2 = \tan(v_1);$ $\delta v_3 = \delta \gamma - \delta v_2;$ $v_{3} = \gamma - v_{2};$ $v_3 = \gamma - v_2;$ $\delta v_4 = \delta \nu * v_2 + \nu * \delta v_2$ $v_4 = \nu * v_2$: $v_4 = \nu * v_2$; $\delta x = \delta v_4 / v_3 + v_4 \delta v_3 / v_3^2;$ $x = v_4/v_3;$ $x = v_4 / v_3;$ $\delta y = \delta \gamma * x + \gamma * \delta x;$ $y = \gamma * x;$ $y = \gamma * x;$

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- The AD code has new input and output variables
 δt, δγ, δν, and δω are new inputs
 δx, δy are new results
- The user must set the input derivatives
 - $\delta t = dt/dp$, $\delta \gamma = d\gamma/dp$, $\delta v = dv/dp$, and $\delta \omega = d\omega/dp$, where p is the parameter to differentiate to
- Examples:
 - Setting $\delta t = 1$, $\delta \gamma = 0$, $\delta v = 0$, and $\delta \omega = 0$, the AD code computes dx/dt and dy/dt
 - Setting $\delta t = 0$, $\delta \gamma = 1$, $\delta v = 0$, and $\delta \omega = 0$, the AD code computes $dx/d\gamma$ and $dy/d\gamma$, etc.
- To get all eight derivatives, the code must be run four times: this is the scalar forward mode





- We can also transform the derivative variables into vectors
 - Using 4-vectors we can compute all derivatives at once
- Example
 - Set $\delta t = [1,0,0,0]$, $\delta \gamma = [0,1,0,0]$, $\delta v = [0,0,1,0]$, and $\delta \omega = [0,0,0,1]$
 - As the result we obtain the full Jacobian matrix $J = \mathrm{D}F$

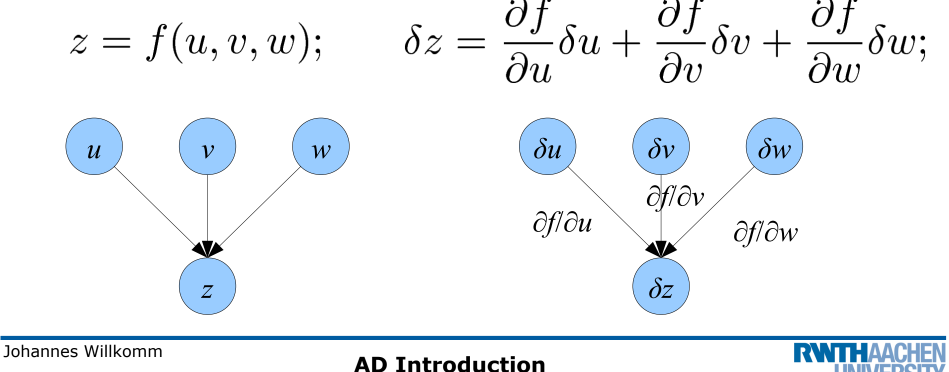
$$\delta x = \left[\frac{\mathrm{d}x}{\mathrm{d}t}, \frac{\mathrm{d}x}{\mathrm{d}\gamma}, \frac{\mathrm{d}x}{\mathrm{d}\nu}, \frac{\mathrm{d}x}{\mathrm{d}\omega}\right]$$
$$\delta y = \left[\frac{\mathrm{d}y}{\mathrm{d}t}, \frac{\mathrm{d}y}{\mathrm{d}\gamma}, \frac{\mathrm{d}y}{\mathrm{d}\nu}, \frac{\mathrm{d}y}{\mathrm{d}\omega}\right]$$

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- To differentiate a program
 - Create new variable δv for each program variable v
 - Differentiate each statement and insert it before the original statement
 - Each δv holds the derivative dv/dp of v w.r.t. the input parameter p



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- AD is also possible by running the program backwards
- For each statement we propagate the derivative of the LHS to the derivatives of the variables on the RHS
 - Create the so-called adjoint statements

$$\begin{split} \delta u &= \delta u + \frac{\partial f}{\partial u} \delta z; \\ \delta v &= \delta u + \frac{\partial f}{\partial v} \delta z; \\ \delta w &= \delta u + \frac{\partial f}{\partial w} \delta z; \end{split} \qquad \begin{array}{c} \delta u & \delta v & \delta w \\ \partial f / \partial v & \partial f / \partial v & \partial f / \partial w \\ \delta z & \delta z & \delta z \end{split}$$

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- Forward sweep
 - The program is executed, saving all variable values
- Initialize adjoints
 - Initialize all derivative variables δv to zero
- Return sweep
 - Execute the adjoint statements in reverse order
 - Now, at any one time, δv contains the adjoint df/dv of v





Lighthouse in reverse

- Run code
- Zero adjoints
- Run adjoint code

$$v_1 = \omega * t;$$

 $v_2 = \tan(v_1);$
 $v_3 = \gamma - v_2;$
 $v_4 = \nu * v_2;$
 $x = v_4/v_3;$
 $y = \gamma * x;$

$$\begin{split} \delta x &+= \gamma * \delta y; \\ \delta \gamma &= 0; \\ \delta \gamma &= 0; \\ \delta \gamma &= 0; \\ \delta \nu &+= v_2 \delta v_4; \\ \delta \omega &= 0; \\ \delta v_2 &+= \nu \delta v_4; \\ \delta v_1 &= 0; \\ \delta v_2 &+= \delta v_3; \\ \delta v_2 &= 0; \\ \delta v_2 &+= -\delta v_3; \\ \delta v_3 &= 0; \\ \delta v_1 &+= \delta v_2 / \cos^2(v_1); \\ \delta v_4 &= 0; \\ \delta \omega &+= t * \delta v_1; \\ \delta t &+= \omega * \delta v_1; \\ \end{split}$$

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- The adjoint code has new in- and outputs
 δx, δy are new inputs
 δt, δy, δv, and δω are new results
- Values for δx and δy are supplied by the user $\delta x = dx/dr$ and $\delta y = dy/dr$ where r is the result to differentiate
- Example
 - Setting $\delta x = 1$ and $\delta y = 0$, the code computes dx/dt, $dx/d\gamma$, dx/dv, and $dx/d\omega$
 - Setting $\delta x = 0$ and $\delta y = 1$, the code computes dy/dt, $dy/d\gamma$, dy/dv, and $dy/d\omega$
- To get all eight derivatives, the code must be run twice, or with 2-vectors as input adjoints





• Given a function

 $\mathbf{y} = F(\mathbf{x}), \quad F : \mathbb{R}^n \to \mathbb{R}^m$

- First order AD computes the Jacobian

 $J = \mathbf{D}F \in \mathbb{R}^{m \times n}$

- Or products thereof
- AD in forward mode
 - Computes Jacobian times vector or Jacobian time matrix products

 Computes vector times Jacobian or matrix times Jacobian products

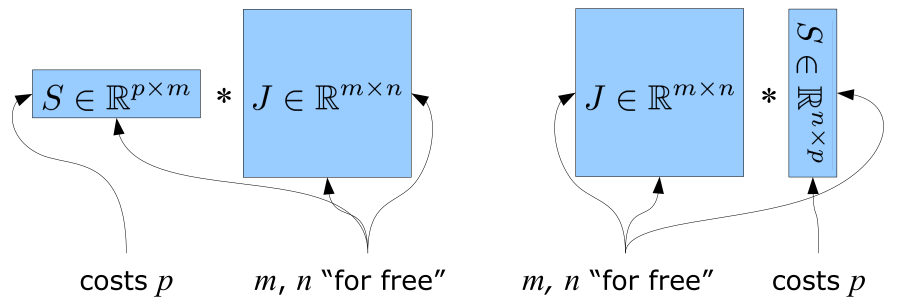
$$J \cdot S, \quad S \in \mathbb{R}^{n \times p}$$

$$S \cdot J, \quad S \in \mathbb{R}^{p \times m}$$





- The time complexity depends on the number of rows or columns in S and the runtime T_F of F
 - Computing J has $T_FO(m)$ in RM and $T_FO(n)$ in FM
 - The c in O is 3 < c < 50, depending on tool & strategy



• Space complexity is $O(T_F)$ in RM!

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- Source transformation
 - New program text is generated
 - Higher order derivatives often not directly supported, but by repeatedly applying the tool
- Operator Overloading
 - Numeric data type (**double**) is replaced by new type
 - Tapeless: Derivatives are stored inside the active variables and updated on the fly
 - Forward mode only
 - With Taping: Computations are first recorded on a so-called Tape, which is then read (forwards or backwards) to compute the derivatives
 - Higher order derivatives are not much more difficult to implement than first order





• Compute polynomial of order *n*

$$F(x, \mathbf{c}) = \prod_{i=0}^{n} c_i x^i$$

- A C-style implementation in MATLAB
 - If x, c_i are all scalars that could also be a one-liner

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Source transformation RM

- Return sweep
 - Zero adjoints
 - Run backwards
 - Compute adjoints

```
• Forward sweep
```

- Run (canonicalized) code
- Save all values overwritten
- Save control flow

```
[a_powX_a_tmpc1] = a_zeros(powX, tmpc1);
 [a_x a_c] = a_z eros(x, c);
if nargin < 3
    [a_r] = a_z \operatorname{eros}(r);
end
[\operatorname{tmpf1}] = \operatorname{pop};
for i=fliplr(1 : tmpf1)
    [powX] = pop;
    a_x = a_x + adjred(x, powX .* a_powX);
    a_{pow}X = adjred(powX, a_{pow}X \cdot \cdot \cdot x);
    [\mathbf{r}] = \operatorname{pop};
    a_tmpc1 = a_tmpc1 + adjred(tmpc1, a_r);
    a_r = adjred(r, a_r);
    [\operatorname{tmpc1}] = \operatorname{pop};
    a_c(i) = a_c(i) + adjred(c(i), a_tmpc1 .* powX);
    a_powX = a_powX + adjred(powX, c(i) .* a_tmpc1);
    [a_tmpc1] = a_zeros(tmpc1);
end
```





AD tools

| ΤοοΙ | Language |
|----------|---------------|
| ADOL-C | C/C++ |
| CppAD | C/C++ |
| ADiFor | Fortran 77 |
| Tapenade | Fortran 77, |
| | Fortran 90/95 |
| ADiMat | Matlab |
| MAD | Matlab |
| ADiCape | CapeML |





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 $\overline{\mathrm{d}}x_k$

- Divided differences
 - Very inaccurate

$$\frac{\mathrm{d}f}{\mathrm{d}x_i} \approx \frac{f(\mathbf{x} + h\mathbf{e}_i) - f(\mathbf{x} - h\mathbf{e}_i)}{2h}$$

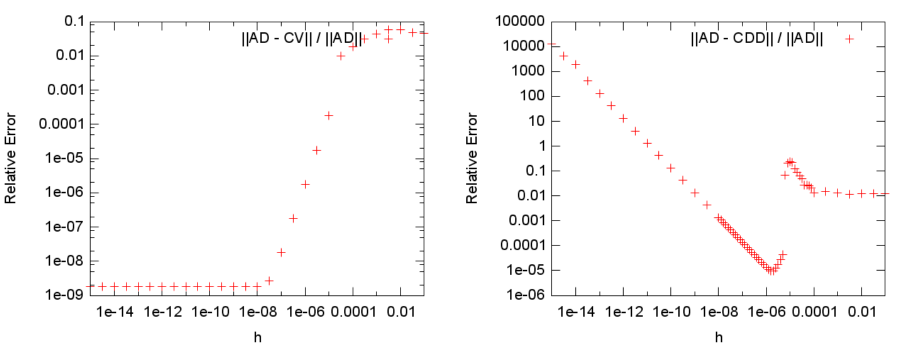
 $\frac{\mathrm{d}f}{\mathrm{d}x_{k}} = \frac{\Im\{f(\mathbf{x} + h\mathbf{i}\mathbf{e}_{k})\}}{h}$

- Difficult to find the right value for h
- \checkmark Only function F is required
- Only Jv with complexity O(n)
- Complex variable method
 - Program needs to be changed similar to AD with OO
 - \checkmark Derivatives are exact, if h is just small enough
 - Need to provide new operations >, <, abs
 - Only Jv with complexity O(n)





CV-Method vs. DD



- The CV-Method is more precise
 - Usually up to machine precision
- And it is safer to use
 - Just set h to a very small value, e.g. $h = 10^{-60}$



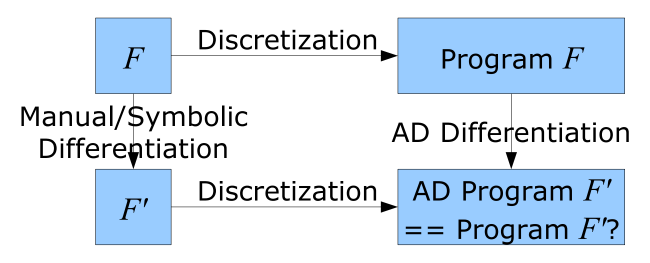


- Symbolic differentiation
 - May be difficult to write a whole program as one expression
 - Large derivative expressions with lots of repeated subexpressions
 - Often very large runtimes
 - Especially for higher order derivatives
 - Differentiation has to be done only once however
- Manual differentiation
 - Usually efficient derivative code
 - Often tedious and error-prone, especially when ${\cal F}$ is changed
 - Discretization of F and F^\prime has to be taken into account





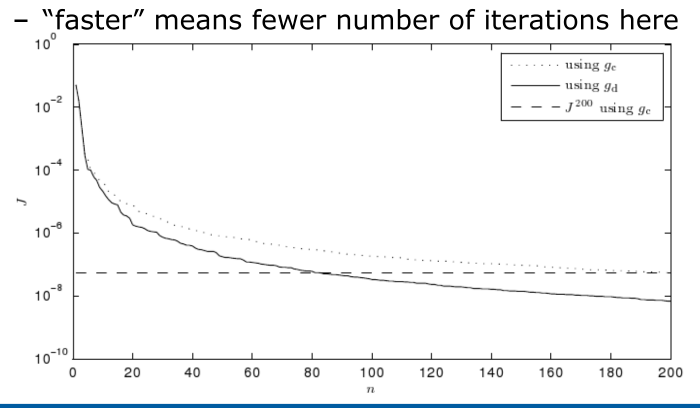
- Let *F* be defined by a PDE
 - Usually implemented by discretization
 - e.g. using the Finite Element Method
- Derivative F' often by discretizing the adjoint PDE
 - The discretization introduces errors in both F and F'
 - AD of the discretized ${\cal F}$ differentiates through the discretization errors of ${\cal F}$







- Solving Inverse Heat Conduction Problem with Conjugate-Gradient optimization using both AD gradient and gradient obtained from adjoint PDE
 - The objective function $J \ensuremath{\operatorname{drops}}$ faster with AD



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- AD advantages
 - AD can provide derivatives of that are efficient, precise, and reliable
 - AD is often easy to apply
- AD disadvantages
 - AD tools can be difficult to use and may lack support for language elements and/or higher order derivatives
 - Applying the reverse mode of AD needs special measures to cope with the memory requirements
 - Possible, but not discussed here
- When you need derivatives you should use AD
- You should consult with an AD expert

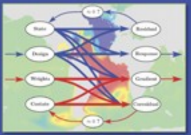




The AD book

"Evaluating Derivatives", 2nd edition Andreas Griewank & Andrea Walther SIAM, Philadelphia 2008 ISBN 978-0-898716-59-7

Andreas Griewank • Andrea Walther



Evaluating Derivatives

Principles and Techniques of Algorithmic Differentiation Second Edition

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