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### Abstract

In optimization problems, derivatives play a crucial role in finding an optimal solution. While automatic symbolic differentiation of matrix expressions and numerical differentiation of tensor expressions are well researched, automatic symbolic differentiation of tensor expressions is less so. In this thesis, we design and implement a tensor calculus that allows for automatic symbolic differentiation of tensor expressions of any order. This calculus uses an Einstein-like product notation that is more general than the notation used in Ricci calculus. [Was sollte hier noch hin?]

## Table of Contents

#### Abstract

#### **Table of Contents**

1	Introduction					
<b>2</b>	A S	A Syntax for Tensor Expressions				
	2.1	Aims	5			
	2.2	Representing Tensor Products	5			
	2.3	Grammar	6			
3	Par	sing and Representation	8			
	3.1	Parsing	8			
		3.1.1 Tokenization	8			
		3.1.2 Tree Building	9			
	3.2	Preprocessing	10			
		3.2.1 Equivalent Subtree Substitution	10			
		3.2.2 Common Subtree Elimination	11			
		3.2.3 Tensor Order Determination	12			
		3.2.4 Axis Propagation	13			
4	Diff	Gerentiation	16			
	4.1	Background	16			
	4.2	Differentiation Rules	17			
		4.2.1 Sum Rule	17			
		4.2.2 Product Rule	18			
		4.2.3 Function Rules	18			
	4.3	First Example	20			
	4.4	Implementation	20			
	4.5	Second Example	23			
	4.6	The Problematic Case of Disappearing Indices	25			
5	${ m Lit}\epsilon$	erature	27			
6	App	pendices	29			
7	7 Declaration of autonomy		30			

### 1 Introduction

With the rise of machine learning applications, interest in optimization problems has been growing. In these kinds of problems, one seeks to find values for a number of parameters that maximize or minimize a given objective function. While some optimization problems have a closed-form solution which can be computed directly, many can only be solved approximately. When computing these approximate solutions, derivatives play a crucial role. In particular, the first derivative of the objective function is used in the ubiquitous gradient descent procedure. In most optimization problems, multiple parameters need to be optimized simultaneously. This means that derivatives need to be computed with respect to multiple variables, organized either in vectors, matrices or higher-order tensors. Commonly used machine learning frameworks (like Tensorflow, PyTorch or Theano) focus heavily on the first derivatives of the objective function.

Higher-order derivatives can, however, also be of interest. As an example, the matrix of second-order partial derivatives, called the Hessian, is used in the application of Newton's method, which can be more efficient than gradient descent for some problems. Convexity checks constitute another application of higher-order derivatives. The convexity of objective functions and their associated optimization problems play a major role in the field of optimization. If a problem is strictly convex or concave, then it has a unique global optimum and the gradient descent algorithm can approximate it arbitrarily well. To check for convexity, the objective function's Hessian matrix needs to be computed and checked for semidefiniteness. If the function's inputs are naturally organized in a matrix or higher-order tensor, then the resulting Hessian will also be a higher-order tensor. The prevalent machine learning frameworks have no way to directly compute these derivatives, and classical computer algebra systems struggle with efficiency, as they work on the level of individual tensor entries.

To bridge this gap, we design and implement a tensor calculus that allows for automatic symbolic differentiation of tensor expressions of any order. Additionally, this calculus will use an Einstein-like notation to represent tensor products, rather than the more complex notation used in Ricci calculus.

This thesis builds on the work by Laue, Mitterreiter and Giesen, who have built a similar calculus for matrix derivatives [2] and developed the theoretical foundation for an Einstein notation based tensor calculus [1].

The structure of the thesis is as follows. First, we develop the syntax of the calculus in form of a grammar. Then, we discuss the parsing and representation of tensor expressions. Following that, we describe the theoretical background and practical implementation of the differentiation process.

## 2 A Syntax for Tensor Expressions

#### 2.1 Aims

In this chapter we develop the language that will be used to specify tensor expressions for differentiation. To differentiate a tensor expression, three pieces of information are needed: 1. The tensor expression itself, 2. The variable with respect to which should be differentiated, 3. The tensor order of all variables in the expression. The neccessity of the first two is obvious, while the third might not immediately be. The orders of each part of the tensor expression need to be known because they play a crucial role in the differentiation rules that will later be implemented.

Our aims when developing the syntax are the following. The syntax should allow representation of tensor expressions of arbitrary order. It should be as general as possible, not focusing on one single interpretation of tensors. Even so, as the motivation for this thesis comes from the background of optimization and machine learning, the most common specifics of these fields should be covered.

The main point of interest for this syntax is how to represent tensors and their products, as there are many ways to multiply two tensors. We start with an established tensor calculus and its notation, and generalize it to fit our problem domain.

### 2.2 Representing Tensor Products

There exists a calculus for tensor expressions, called Ricci calculus, that is heavily used in physics. The notation it uses allows for tensors to represent both n-dimensional arrays and multilinear functions. The distinction between the two forms occurs by the location of their indices, either in sub- or superscript. For example, the inner product of two vectors,  $y^T x$ , would be written as  $y_i x^i$ . And the Matrix-vector product Ax would be  $A_j^i x^j$ . While Ricci calculus can be useful, for example to physicists, the many indices on each tensor make it less readable.

Since, for our purposes, tensors are simply containers for variables, we have no need to expressly define some tensors as multilinear functions. A part of Ricci calculus that we bring over into our notation is a derived version of the Einstein summation convention. This is a notational convention that allows elegant representation of tensor products. Since tensors, depending on their order, may be multiplied in many ways, we need a way to describe which tensor axes should be multiplied in what way. For example, two vectors could be multiplied by taking their inner product, their outer product, or even their elementwise product. Einstein notation handles

this problem by adding index strings to the inputs and the output of the product, each index representing an axis in the respective tensor. If an index appears in both inputs, the respective axes will be multiplied. If an index appears in inputs but not in the output, this axis will be summed over. In true Einstein-notation, output indices are omitted, so indices that appear in both inputs are always summed over. For more generality, we include output indices. This simple set of rules provides an elegant way to express the main tensor operations.

We take a large part of the syntax for tensor multiplication from the established Python software package Numpy, specifically its einsum procedure. A product of two tensors x and y has the form  $x *_{(a,b\to c)} y$ , where a,b and c are strings of indices. A formal characterization of the product  $C = A *_{(s_1,s_2\to s_3)} B$  is given in Equation 2.1.

$$C[s_3] = \sum_{(s_1 \cup s_2) \setminus s_3} A[s_1] \cdot B[s_2]$$
 (2.1)

As an example of tensor products, see the inner, outer and elementwise product of two order 1 tensors (vectors) x and y:  $x *_{(i,i \to)} y$ ,  $x *_{(i,j \to ij)} y$ ,  $x *_{(i,i \to)} y$ . The application of a matrix A to a vector v would be written as  $A *_{(ij,j \to i)} v$ , while the elementwise product of two matrices A and B would be  $A *_{(ij,ij \to ij)} B$ . By swapping the order of output indices, the axis order can be manipulated. A special case of this is the matrix transpose, which for a matrix X can be taken using the product  $X *_{(ij,\to ji)} 1$ . One may even take diagonals of tensors using this notation, the inner product of a square matrix A's main diagonal and a vector v would be  $A *_{(ii,i\to)} v$ .

### 2.3 Grammar

With the representation of tensor products handled, we can now define the full syntax grammar. The grammar, presented here in extended backus-naur form, is based on a standard grammar for mathematical expressions by Julien Klaus. It was adapted to tensor expressions by generalizing multiplication to tensor products using the einstein summation convention.

Here  $\{x\}$  describes that x may occur any number of times, while  $\{x\}$ + lets x appear at least once.

As can be seen in the grammar, we allow the following binary operations on tensors: products (\*(,->)), sums (+), differences (-) and quotients (/). With the exception of tensor products, which have already been discussed, these operations are executed elementwise. Exponentiation (^) is also included, but not as a true binary operator on tensors, as the exponent may only be scalar, with each entry of the basis tensor being combined with the exponent. This is not apparent from the grammar, as tensor variable order is specified separately from the tensor expression itself.

A variety of functions with relevance to optimization and machine learning are included, mostly being applied elementwise to tensors of any order. The exceptions to this are the matrix inverse <code>inv</code>, the matrix determinant <code>det</code> and the adjugate matrix <code>adj</code>, all of which are not applied elementwise and may only operate on order 2 tensors. The inclusion of these functions takes away from the generality of our grammar, as the idea of an inverse and a determinant can only be applied to matrices, not higher-order tensors, but they are crucial to some common optimization problems that work with matrices.

## 3 Parsing and Representation

Having defined the syntax, we now discuss how a given tensor expression is parsed and represented internally. In our application, tensor expressions are represented using directed acyclic graphs derived from binary trees. This data structure was chosen because graphical representations are well tested for expressions and because the foundational paper by Laue, Mitterreiter and Giesen also assumes such a representation. This allows the differentiation rules stated in the paper to be easily implemented. Furthermore, many of the algorithms described in this chapter and the next are most naturally written recursively. A recursive data structure like a tree lends itself to these algorithms. In this chapter we first describe how a tensor expression is parsed into a binary expression tree, then explain the processing steps taken to transform the binary tree into a directed acyclic graph without duplicate subexpressions, and also describe how we assign a tensor order and an axis tuple to each node in the graph.

### 3.1 Parsing

The process of turning a string into a different data structure that allows for further processing is called parsing. It is split into two steps, tokenization and tree building.

#### 3.1.1 Tokenization

To be able to parse an expression string, we first transform it into a sequence of tokens. This is done by the tokenizer, sometimes also called lexical scanner, which recognizes certain defined patterns in the expression string and unifies them into a token. Each of these tokens represents a logical unit that is used in the next step of parsing. Tokenization allows the parser to work only with well-defined tokens, not having to work on the level of individual characters. Each token is made up of two parts, the descriptor and the identifier. The descriptor is one of a few defined categories, describing what kind of logical unit the token forms, while the identifier contains the characters the token was formed from. As an example, the string A + sin(b) would be tokenized into the following sequence: (ALPHANUM, 'a'), (PLUS, '+'), (ELEMENTWISE\_FUNCTION, 'sin'), (LRBRACKET, '('), (LOWERCASE\_ALPHA, 'b'), (RRBRACKET, ')').

We define token descriptors for constants (e.g. '1.05'), variable names (e.g. 'A', 'b', 'C1'), special symbols ('+', '-', '\*', '/', '^', ', ', '(', ')', '>'), keywords ('declare', 'expression', 'derivative', 'wrt') and function names, which are recognized using the list of functions given in the grammar (e.g. 'sin', 'abs',

'det'). There are two token descriptors for functions, ELEMENTWISE\_FUNCTION and SPECIAL\_FUNCTION, because derivative rules are different for these two types. Special functions are the matrix determinant det, the matrix inverse inv and the adjugate matrix adj.

#### 3.1.2 Tree Building

The output of the tokenizer is used to build a binary tree representing the expression. A binary tree is a type of tree where each node has at most two child nodes. Like in other trees, each node has exactly one parent node. The exception to this is the root, which has no parent. We call nodes that have no children leaf nodes.

Formally, we define a binary tree B as follows: B is either the empty set or a tuple (left(B), root(B), right(B)), where left(B) and right(B) are also binary trees and have B as their only parent, and root(B) is the root of the tree, containing some value.

The process of constructing such a tree from a given expression follows the structure of the grammar defined in the previous chapter. Starting with expr, we recursively step through the grammar, using the tokens to decide which options to follow. For each of the words defined in the grammar, there is a function in our parser. When a certain word is expected, the function is called. For example, at the start, expr is called, which first calls term, which calls factor, which then looks for any MINUS-tokens before calling atom, which then checks if there is a constant, a function application, a variable name, or another expression, and so forth. Thus, we traverse the expression from left to right while building binary tree nodes when appropriate. Leaf nodes are created when we reach constants or variable names, and they are connected using branch nodes containing operators or functions. Each node, with its subtree, represents a subexpression, with the root node of the whole tree representing the entire expression. If there is ever a point at which expectations are not met, for example, if a left round bracket token ((LRBRACKET, '('))) is expected but not found, we abort program execution.

As an example, we take the expression from before, A + sin(b), tokenized to (ALPHANUM, 'a'), (PLUS, '+'), (ELEMENTWISE\_FUNCTION, 'sin'), (LRBRACKET, '('), (LOWERCASE\_ALPHA, 'b'), (RRBRACKET, ')').

The order of function calls generated when parsing this expression would be the following:

- expr
- term
- factor
- atom We identify 'a' as a variable name, generate a leaf node and jump back up the call stack to expr. There, we find the expected '+' and add a sum node as the parent of our leaf node.
- term
- factor

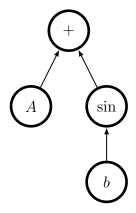


Figure 3.1: Binary expression tree generated from the expression string A + sin(b). Empty nodes are omitted.

- atom We identify 'sin' as a function name and generate an elementwise-function node as the second child of our sum node. We also find the expected '('.
- expr
- term
- factor
- atom We identify 'b' as a variable name and generate another leaf node as the child of our elementwise-function node. We jump back up the call stack, find the expected ')', jump all the way to the top of the call stack and finish.

The binary tree generated from this can be seen in Figure 3.1. Note that, by convention, we set the argument of a unary function node to its right child.

### 3.2 Preprocessing

Having shown how a binary expression tree is created from a given tensor expression, we now describe how this tree is processed further to prepare it for differentiation. We employ the following three preprocessing steps.

- Some subtrees are substituted with equivalent ones which are more easily differentiated.
- Subtrees which occur more than once are fused.
- Each node's output tensor order is determined and stored.

### 3.2.1 Equivalent Subtree Substitution

Some operations or functions may be viewed as equivalent to, or as a special case of another, more easily differentiable operation or function. In these cases, we substitute the more general case to reduce the number of differentiation rules that

need to be implemented. For example, when encountering the expression A - B, we substitute A + -(B), viewing (-) as an elementwise function. This allows us to use the differentiation rules for sums and elementwise functions, not having to implement a new rule for differences. A full overview of the substituted expressions can be seen in Table 3.1.

Original	Substitute
A-B	A + -(B)
$A^B$	$\exp(B *_{(,->)} \log(A))$ A.* elementwise_inverse(B)
A/B	$A$ .* elementwise_inverse( $B$ )
adj(X)	$\det(X) *_{(ij \to ij)} \operatorname{inv}(X)$

Table 3.1: Table showing all operations that get substituted during preprocessing, along with their substitutes. Note that (.\*) is a placeholder for elementwise multiplication. It is replaced with the appropriate tensor product during tensor order determination. elementwise\_inverse is an elementwise function for which a differentiation rule is implemented.

#### 3.2.2 Common Subtree Elimination

When differentiating, we need to be able to tell when a subexpression is reached multiple times. To help with this, and also to reduce the number of nodes in the expression tree, we fuse subtrees that occur more than once.

First, to identify duplicate subtrees, the notion of equality needs to be defined for trees. We define equality of two trees, A and B, as follows.

$$A = B \iff \operatorname{root}(A) = \operatorname{root}(B) \land \operatorname{left}(A) = \operatorname{left}(B) \land \operatorname{right}(A) = \operatorname{right}(B)$$

Equality of nodes is handled by comparing all components of the node (node type, name, indices for product nodes) individually.

Now that we may identify when two subtrees are equal, we continue with common subtree elimination. We first obtain all subtrees by traversing the whole tree. Then, we store each unique subtree we find in a hash table. When we encounter a subtree that is already present in our hashmap, we let its parent node have the tree in the hashmap as its child instead of the original. In doing so, we remove duplicate subtrees and transform the binary expression tree into a directed acyclic graph (DAG) representing the expression. Nodes in the DAG still only have two child nodes, but may have many parent nodes.

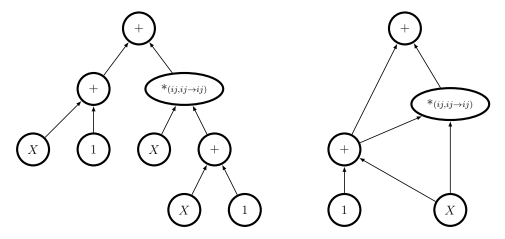


Figure 3.2: Left: Binary tree generated representing the expression  $(X + 1) + X *_{(ij,ij\to ij)} (X + 1)$ . Right: DAG obtained by eliminating common subtrees in the left tree.

#### 3.2.3 Tensor Order Determination

Another preprocessing step is tensor order determination. During differentiation, we often create new product nodes. Because we need to specify product indices when we create a product node, we require knowledge of the tensor orders of both inputs and the output of the new node, which depend on the tensor orders of nodes in the original DAG. Because we may need the tensor rank of all nodes in the original DAG, we determine them by propagating tensor orders from the bottom of the DAG upwards toward the root.

The bottom nodes contain either variables or constants. The tensor orders of variables need to be given, as they function as the start for the upwards propagation of tensor orders. Each parent node calculates its tensor order based on its child nodes. For example, a sum will have its order set equal to the order of its children. A full overview of these order determination rules can be found in table 3.2.

So far we have not addressed how the tensor order of a constant is determined. Constants may need to be broadcast to fit into the expression. For example, in the expression  $1 + \det(X)$  with an order 2 tensor X, 1 would be a scalar (order 0 tensor) since it needs to be summed with another scalar. But in the expressio  $\det(X+1)$ , 1 would need to be an order 2 tensor filled with ones, because it is summed with another order 2 tensor. To accomplish this broadcasting, constants have their order set to -1 to start with. Then, when an error is encountered during tensor order determination because an order -1 constant does not fit into a part of the expression, the order of the constant is set to the value that makes it fit.

In a sum, this would be the order of the other summand. In a product, the order would be determined by the given input indices. In a special function, the order is the input order the function operates on. Since all our special functions operate on matrices, this is always 2. The case of an elementwise function is more difficult, since they operate on arbitrary input tensor orders. Nonetheless, we may find a fitting order using the expression in which the output of the elementwise function is used, by using the order that is needed there. This is possible since the input and output order of elementwise functions are always the same. As an example, the order of the

tensor constant 2 in the expression  $X + \log(2)$  may be determined by finding out what order  $\log(2)$  needs to be to fit in the sum. Since summands always have the same order,  $\log(2)$ , and by extension 2, need to have the same order as X. Using these rules, we determine the tensor order of any node in an expression DAG.

Node Type	Tensor order
SUM	$\operatorname{order}(\operatorname{left}(X))$
PRODUCT	$output\_indices(X)$
ELEMENTWISE FUNCTION	$\operatorname{order}(\operatorname{right}(X))$
adj, inv	2
det	0

Table 3.2: Rules applied to determine tensor orders of a node X, depending on the type of node it is. Note that the output\_indices of a product node are given after the  $\rightarrow$  in the product string, that the sum may also take its order from the right child which has the same order, and that each special function (det, inv, adj) needs its own rule.

### 3.2.4 Axis Propagation

In addition to the tensor order, we also require each node to store what its axes are. The general structure of tensor derivatives is invariant to the specific axis lengths of the tensors variables. For example, the derivative of  $X *_{(ij,ij\to)} X$  with respect to the order 2 tensor X is always X + X or  $2 *_{(,ij\to ij)} X$ , no matter whether  $X \in \mathbb{R}^{3\times 5}$  or  $X \in \mathbb{R}^{400\times 2}$ . This is why we do not require the length of each axis of each variable in a tensor expression to differentiate it.

However, knowing how the shape of each node depends on the shape of the input variables is still useful. It is necessary for generating program code to numerically compute the derivative we determine and gives us more information about the structure of the derivative. For these reasons, we do not only propagate tensor order through the input and differentiation DAGs, but also the axes.

First, we give each node a tuple of axes. Note that we simply use integers to refer to axes in our DAGs. A node X with tensor order 2 may for example have the axes (1,2), where 1 and 2 are simply names for axes, not containing any further information. In the same way we propagated tensor order, we start from the leaf nodes of the DAG and move toward the top.

In the leaf nodes of the DAG are variables and constants. Constants start out with no axes while they still have the initial tensor order -1. When a constant gets broadcast, it is assigned fitting axes depending on its surroundings in the expression, in the same way they get fitting tensor orders.

Variables are assigned an n-tuple of axes, n being the tensor order of the variable. Each axis is newly created and not used by any other variable at first. Then, these axes are propagated upwards through the DAG, using similar rules to the propagation of tensor orders, visible in table 3.3.

The rule for products is slightly more complex. It chooses the correct axes for the product node from the axes of its children based on the product index strings. See Algorithm 1 for further details.

Node Type	Axes
SUM	axes(left(X))
PRODUCT	See Algorithm 1
ELEMENTWISE FUNCTION	axes(right(X))
adj, inv	axes(right(X))
det	()

Table 3.3: Rules applied to determine axes of a node X, depending on the type of node it is.

```
Algorithm 1 Propagate_Product_Axes_Bottom_Up (node)

for each i in output_indices(node) do

if i is in left_indices(node) then

p \leftarrow \text{position of } i in left_indices(node)

append axes(left(node))[p] to axes(node)

else if i is in right_indices(node) then

p \leftarrow \text{position of } i in right_indices(node)

append axes(right(node))[p] to axes(node)

end if

end for
```

After this procedure, each node has a fitting tuple of axes. However, many of these axes with different names are actually the same. For example, in the expression A + B for two order 2 tensors A and B, this procedure may assign A the axis tuple (1,2) and B the axis tuple (3,4). A and B are assigned different axes even though the sum requires axis 1 to be equal to axis 3 and axis 2 to be equal to axis 4.

This is why we pass through the DAG a second time, going from the top toward the bottom, unifying axes that are the same. For each node x, we set its childrens' axes depending on its own, with different procedures for different node types.

Node Type	Left Child Axes	Right Child Axes
SUM	axes(X)	axes(X)
PRODUCT	See Algorithm 2	See Algorithm 2
ELEMENTWISE FUNCTION	-	axes(X)
adj, inv	-	axes(X)
det	-	()

Table 3.4: Rules applied when propagating axis downward through a DAG.

An overview of the applied propagation rules is seen in Table 3.4. For a product nodes we employ a more complex procedure, setting each child's axes to the node's axes depending on the product indices. More details may be seen in Algorithm 2.

After this second pass through the DAG, each node now contains a fitting tuple of axes, unified as much as possible, with each axis coming from a variable in the original DAG.

#### Algorithm 2 Propagate\_Product\_Axes\_Top\_Down (node)

```
for each i in output_indices(node) do
    p \leftarrow \text{position of } i \text{ in output\_indices}(node)
    for each j in left_indices(node) do
         q \leftarrow \text{position of } j \text{ in left\_indices}(node)
         if i = j then
             axes(left(node))[q] \leftarrow axes(node)[p]
         end if
    end for
    for each j in right_indices(node) do
         q \leftarrow \text{position of } j \text{ in right\_indices}(node)
         if i = j then
             axes(right(node))[q] \leftarrow axes(node)[p]
         end if
    end for
end for
for each i in left_indices(node) do \triangleright Unifies equivalent left and right child axes
    p \leftarrow \text{position of first occurrence of } i \text{ in left\_indices}(node)
    if i exists in right_indices(node) then
         q \leftarrow \text{position of first occurrence of } i \text{ in right\_indices}(node)
         l \leftarrow \operatorname{axes}(\operatorname{left}(node))[p]
         r \leftarrow \operatorname{axes}(\operatorname{right}(node))[q]
         rename axis l to r in all nodes of the DAG
    end if
end for
```

### 4 Differentiation

The previous chapter was concerned with creating a suitable internal representation for tensor expressions, which turned out to be a directed acyclic graph (DAG). In this chapter, we use this DAG representation of the expression for the differentiation. We start by giving some background information on the general differentiation procedure and describe the differentiation rules in theory. Following that, we elaborate on the details of our implementation of the procedure and the rules.

### 4.1 Background

The theory for differentiating tensor expressions using our Einstein-like notation was developed by Laue, Mitterreiter and Giesen. They developed differentiation rules for DAG representations of tensor expressions based on the Fréchet Derivative. In their paper, they present two modes of differentiation, forward and reverse mode. This thesis utilizes reverse mode.

In reverse mode, the computation traverses the DAG from top to bottom. The goal is to calculate for each node v the derivative of the whole expression y with respect to v. This value,  $\frac{dy}{dv}$ , is also called the pullback of v. When traversing the DAG, the pullback is known for each parent node u of v, so we may apply the chain rule:

$$\frac{dy}{dv} = \sum_{u:(v,u)\in E} \frac{dy}{du} \cdot \frac{du}{dv}$$
(4.1)

where  $(v, u) \in E$  means that there is an edge going from v to u in the DAG, which makes u a parent of v. Note also that  $\cdot$  is a placeholder for an appropriate tensor product, which is specified in the differentiation rules. This rule allows us to calculate the pullback of v using the known pullbacks of v's parents u. The only new calculation to be done for each node is the derivation of the parent nodes u with respect to v,  $\frac{du}{dv}$ .

The procedure starts with the topmost node y, representing the entire expression, and computes its pullback, which is the derivative with respect to itself. This pullback is then used to compute the pullback of y's child nodes, which are used to compute their child nodes, continuing recursively until all paths from the top node y to the argument node have been traversed. The pullback for the node containing the argument is the result of the differentiation, the derivative of the whole expression y with respect to the argument.

$$\begin{pmatrix} x_1 \\ x_2 \\ x_3 \end{pmatrix} \qquad \begin{pmatrix} \frac{\partial x_1}{\partial x_1} & \frac{\partial x_2}{\partial x_1} & \frac{\partial x_3}{\partial x_1} \\ \frac{\partial x_1}{\partial x_2} & \frac{\partial x_2}{\partial x_2} & \frac{\partial x_3}{\partial x_2} \\ \frac{\partial x_1}{\partial x_3} & \frac{\partial x_2}{\partial x_3} & \frac{\partial x_3}{\partial x_3} \end{pmatrix} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}$$

Figure 4.1: Left: An order 1 tensor. Right: The derivative of the left tensor with respect to itself, resulting in an order 2 delta tensor.

In the very first step of the procedure, the derivative of the top node y with respect to itself is computed. Since y is a tensor-valued function, its derivative is also a tensor. Specifically, differentiating y with respect to itself means that each entry of the tensor represented by y is differentiated once with respect to all other entries. Thus, this differentiation creates a new tensor of the same size as y for each entry in y, where for each of these tensors, only one entry is 1, while the rest are 0, visible in Figure 4.1. Concretely, if y is an order n tensor, then its derivative is a tensor  $\delta_n$  of order 2n, where

$$\delta_n[i_1, ..., i_{2n}] = \begin{cases} 1 & \text{if } (i_1, ..., i_n) = (i_{n+1} ... i_{2n}) \\ 0 & \text{else} \end{cases}$$
 (4.2)

Intuitively, the first n indices point to the entry of y that is being differentiated, while the last n indices point to the entry with respect to which we are differentiating. Only when we differentiate an entry with respect to itself is the result a 1, otherwise it is 0. Tensors that satisfy Equation 4.2 are called delta tensors. Note that we call a delta tensor of order 2n by the name  $\delta_n$  and not  $\delta_{2n}$ , because delta tensors may only have even order. The first node created in the differentiation procedure is always a delta tensor representing the topmost node y differentiated with respect to itself.

### 4.2 Differentiation Rules

Having established the basic differentiation procedure, we will now describe the rules used to compute the derivative  $\frac{du}{dv}$  for a node v and its parent u. Proofs for the correctness of these rules can be found in the paper [1] by Laue, Mitterreiter and Giesen. Since we are computing  $\frac{du}{dv}$ , the applied rule depends on the type of node u is.

#### 4.2.1 Sum Rule

If u is a sum node, the differentiation is very simple. In this case, u adds nothing of its own to the pullback of v, so the contribution of u to the pullback of v is only  $\frac{dy}{du}$ . Thus, when differentiating a sum node that is their childrens only parent, like the one seen in Figure 4.2, the pullback of the children is simply the pullback of the sum node.

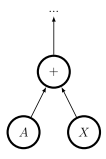


Figure 4.2: A sum node in an expression DAG. The pullbacks of its children A and X are the same as its pullback.



Figure 4.3: **Left**: A product node representing a scalar-vector-product in an expression DAG. "..." represents the rest of the DAG. **Right**: Derivative of the left DAG with respect to x computed using the product rule. "..." represents the pullback of the product node. Note that the product indices in the right DAG depend on the indices s4 of the topmost node in the left DAG (not visible here), which is assumed to be scalar in this case.

#### 4.2.2 Product Rule

If u is a product node representing the subexpression  $v *_{(s_1,s_2 \to s_3)} w$  with index strings  $s_1$ ,  $s_2$  and  $s_3$  in a tensor expression where the topmost node has indices  $s_4$ , then we may apply the product rule as follows. The contribution of u to the pullback of v is

$$\frac{dy}{du} *_{(s_4s_3, s_2 \to s_4s_1)} w,$$

and the contribution to the pullback of w is

$$\frac{dy}{du} *_{(s_4s_3, s_1 \to s_4s_2)} v.$$

An example of how the product rule is applied to an expression DAG can be seen in Figure 4.3. There is a caveat to the product rule, because by itself it is not applicable to all tensor products. This detail will be discussed further in the section on implementation details.

#### 4.2.3 Function Rules

If u is a function node, we distinguish between two cases. The node u either represents an elementwise function or a special function. The elementwise case allows for a slight simplification over the special function case.

If the node u represents an elementwise function f being applied to the child node v, then the contribution of u to the pullback of v is

$$\frac{dy}{du} *_{(s_2s_1, s_1 \to s_2s_1)} f'(v),$$

where  $s_1$  is the index string of v,  $s_2$  is the index string of the output node of the DAG and f' is the derivative of f. An example can be seen in Figure 4.4.

If the node u represents a special function f being applied to the child node v, then the contribution of u to the pullback v is

$$\frac{dy}{du} *_{(s_3s_2, s_2s_1 \to s_3s_1)} f'(v),$$

where  $s_1$  is v's input, representing the domain of f,  $s_2$  is the index string of v, representing the range of f, and  $s_3$  is the index string of the output node of the DAG with f' still being the derivative of f. The distinction between  $s_1$  and  $s_2$  is neccessary because domain and range of f may not be the same with these functions.

The derivatives f' of all accepted functions f need to be known. They are listed in Table 4.1

Function	Derivative wrt. x
$\overline{-(x)}$	-(1)
$\sin(x)$	$\cos(x)$
$\cos(x)$	$-(\sin(x))$
$\tan(x)$	$1./(\cos(x).*\cos(x))$
$\arcsin(x)$	$1./(1+-(x.*x))^{0.5}$
$\arccos(x)$	$-(1./(1+-(x.*x))^{0.5})$
$\arctan(x)$	1./(1+(x.*x))
$\exp(x)$	$\exp(x)$
$\log(x)$	1./x
tanh(x)	$1 + -(\tanh(x).*\tanh(x))$
abs(x)	$ \operatorname{sign}(x) $
sign(x)	0
relu(x)	$\operatorname{relu}(\operatorname{sign}(x))$
1./x	-(1./(x.*x))
inv(x)	$-(\operatorname{inv}(x)) *_{(ij,kl \to iklj)} \operatorname{inv}(x)$
$\det(x)$	$\mid \operatorname{adj}(x) *_{(ij, \to ji)} 1$

Table 4.1: Derivatives of all used functions. .\* is a placeholder for elementwise multiplication with the fitting indices. The elementwise inverse 1./x is internally handled as an elementwise function, which is why its derivative is listed here.



Figure 4.4: **Left**: Part of a DAG that contains the subexpression  $\sin(X)$ . X is an order 2 tensor (matrix) and "..." represents the rest of the DAG. **Right**: Derivative of the left DAG with respect to X computed using the rule for elementwise functions. "..." represents the pullback of the function node. Note that the topmost output node of the left DAG is assumed to be scalar here, which has an impact on the indices in the product node in the right DAG.

### 4.3 First Example

In this section, we present a detailed example of the differentiation procedure. Consider the simple DAG in Figure 4.5, representing the expression  $X *_{(ij,ij\to ij)} X$ , the elementwise product of a matrix X with itself. We will be differentiating this DAG with respect to X. Each step of the process is visualized in Figure 4.6.

Following the differentiation procedure, we start by differentiating the topmost node with respect to itself. Since the expression is an order-2 tensor, this creates a delta tensor  $\delta_2$  of order 4. (Step 1 in Figure 4.6)

Continuing from the topmost node, we compute its contribution to the pullback of the left child first. In this case, the left child and the right child are the same, but are still handled separately. Using the product rule, we can see that the contribution of the top node y to the left child is  $\frac{dy}{dy} *_{(abij,ij \to abij)} X$ , with  $\frac{dy}{dy}$  being  $\delta_2$ . (Step 2 in Figure 4.6)

Since X has no children, we continue with the right child of y. Here, we apply the product rule and obtain the same  $\delta_2 *_{(abij,ij \to abij)} X$  as the contribution to the pullback of X. As contributions to the pullbacks of the same node need to be added (see Equation 4.1), we add a sum node as the root of our differentiation DAG. (Step 3 in Figure 4.6)

Because all ways from the root node to the argument have been processed, we are finished. The final DAG, simplified using common subtree elimination, can be seen in Figure 4.7.

### 4.4 Implementation

In this chapter we describe how the reverse mode differentiation procedure is implemented using the DAG representation of tensor expressions and the differentiation rules introduced in the previous sections.

An overview of the differentiation procedure can be seen in Algorithm 3 and

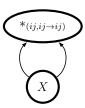


Figure 4.5: Expression DAG representing the elementwise product of a matrix X with itself.

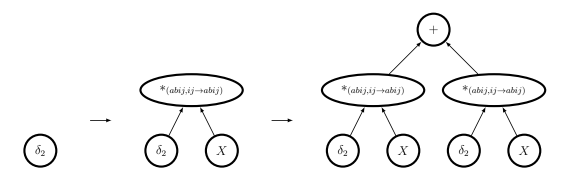


Figure 4.6: Progression of the differentiation procedure on the DAG in Figure 4.5. Left: Pullback of the topmost node. Middle: One contribution to the pullback of X. Right: Pullback of X, both contributions summed.

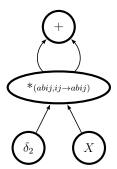


Figure 4.7: Differentiation DAG of the original DAG in Figure 4.5, simplified using common subtree elimination.

#### **Algorithm 3** Differentiate (root, argument)

```
diffdag \leftarrow \text{delta}(\text{order}(root)) \triangleright Creates a node representing \delta_{\text{order}(root)} return ReverseModeDiff(root, argument, diffdag, EmptyDictionary())
```

#### **Algorithm 4** ReverseModeDiff (node, argument, diffdag, contributions)

```
if subtree of child does not contain argument then
Continue with next child
end if
Apply fitting differentiation rule (depending on node) to child
if child is in keys(contributions) then
Add new contribution to contributions[child] using a sum node
Move all parents of the contributions[child] to the sum node
Update contributions[child] as the sum node
else
contributions[child] ← diffdag
diffdag ← ReverseModeDiff(child, argument, diffdag, contributions)
end if
end for
return diffdag
```

As can be seen in Algorithm 3, the first node is added specially, before the real differentiation algorithm starts. This node, a delta tensor, is the derivative of the top node y with respect to itself and has double the order of the original top node.

The original DAG is traversed from the top node down to the argument node, building up the differentiation DAG alongside it. With each node u of the original DAG that is processed, its contributions to the pullbacks of its child nodes are added to the differentiation DAG. These pullbacks are computed using the differentiation rules discussed in the previous chapter.

As we are operating on a DAG and not a tree, a single node may be reached multiple times during differentiation. When differentiating a node u, one of its children v may have already been reached through a different parent. In this case, a part of the pullback of v has already been computed and is present in the differentiation DAG. The new pullback contribution we computed from u needs to be added to the old part, as can be seen from Equation 4.1. To be able to do this, we keep a dictionary mapping from nodes in the original DAG which we have already reached to their pullback in the differentiation DAG. This lets us add the old and new pullback contributions with a sum node. Afterwards, we update the entry in the dictionary to the sum node that was just added so that further contributions are added there. We also need to move all nodes in the differentiation DAG that rely on the pullback of v, specifically the parents. Thus, we set all parents of the old pullback to have the updated pullback as their child. This way, we build up the correct pullback over time, adding contributions as we find new ways to reach each node.

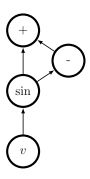


Figure 4.8: DAG representing the expression  $\sin(v) + -(\sin(v))$ .

### 4.5 Second Example

To better illustrate the process of building a pullback piece by piece, we showcase an example. Consider the DAG in Figure 4.8 representing the expression  $\sin(v) + -(\sin(v))$  for an order 1 tensor v. We differentiate this expression with respect to v.

Starting out, we compute the pullback of the topmost node. Since the output of the topmost node is an order 1 tensor, its pullback, the derivative with respect to itself, is an order 2 delta tensor  $\delta_1$ . (Step 1 in Figure 4.9)

Next, we compute the contribution of the topmost node to its children's pullbacks. However, since the topmost node is a sum node, it adds nothing of its own, which makes its childrens pullbacks the same as its.

Continuing with the node's children, we start (arbitrarily) with the left child. The left child is an elementwise function node representing  $\sin(v)$ . Using the differentiation rule for elementwise functions along with the derivative of the sin function, we see that its contribution to its child's pullback is  $\delta_1 *_{(ba,a\to ba)} \cos(v)$ . This is the first contribution to the pullback of v. (Step 2 in Figure 4.9) Since the only child of the sin node is the argument node v, we need not continue further and instead jump back up to explore the second child of the topmost node.

The second child is an elementwise function node representing  $-(\sin(v))$ . Using the differentiation rule for elementwise functions along with the derivative of the (-) function, we get  $\delta_1 *_{(ba,a \to ba)} -1$  as the contribution of the node to its child's pullback.

The node's child is the sin node we have been to before. This means we need to add the new contribution to this node's pullback to the old one. The old contribution to the pullback of  $\sin(v)$  was  $\delta_1$ , to which we now add  $\delta_1*_{(ba,a\to ba)-1}$  using a sum node. Since the pullback of v depends on the pullback of  $\sin(v)$ , we need to make sure to update it. We manage this by moving all parents of the old pullback to be parents of the new pullback instead. The left child of the product  $\delta_1*_{(ba,a\to ba)}\cos(v)$  gets updated to be the sum node in the new, complete pullback. (Step 3 in Figure 4.9)

Since there are no more unvisited children, we are done. The finished differentiation DAG represents the expression  $(\delta_1 + (\delta_1 *_{(ba,a\to ba)} -1)) *_{(ba,a\to ba)} cos(v)$ . The complete, simplified DAG is visible in Figure 4.10.

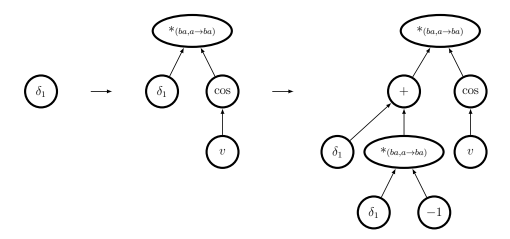


Figure 4.9: Progression of the differentiation procedure on the DAG in Figure 4.8. Left: Pullback of the +, sin and (-) nodes. Middle: First contribution to the pullback of v. Right: Full pullback of v.

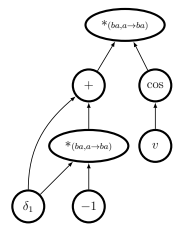


Figure 4.10: Full simplified differentiation DAG of the original DAG in Figure 4.8.

### 4.6 The Problematic Case of Disappearing Indices

In this section we present a problem with the differentiation of tensor products that was discovered during development, along with a solution.

In the proof of the reverse mode differentiation product rule in the paper by Laue, Mitterreiter and Giesen [1] (Theorem 8) in the fourth equality, the associative property for tensor products is used to transform  $\bar{C} *_{(s_4s_3,s_3\to s_4)} (A *_{(s_1,s_2\to s_3)} h)$  into  $(\bar{C} *_{(s_4s_3,s_1\to s_4s_2)} A) *_{(s_4s_2,s_2\to s_4)} h$ . However, as can be seen in the corresponding Lemma 1 on the previous page of the paper,

For some DAGs, this causes invalid product nodes to be created during differentiation. Consider the case where an index appears only in the index string of one input of a product node, but not in the output index string. An example of this would be the DAG on the left in Figure 4.11, containing a node representing the product  $A *_{(ij,j\to)} v$ , where the index i only appears in the left index string. When differentiating this DAG with respect to A, we compute the contribution of the product node to the pullback of A and receive  $\delta_0 *_{(,j\to ij)} v$  using the product rule. The resulting product node is not valid, because the index i appears in the result index string, but not in either input index string.

Even though these cases create invalid products, they can still be interpreted in an intuitive way. In our example, one of A's axes is summed over in the original product. Because of this, the derivative over this axis will be the same for all entries. If we removed this redundancy, we would obtain  $\delta_0 *_{(,j\to j)} v$  as the result. The extra index i in the result indices of the product in the differentiation DAG can be interpreted as duplicating the result over another axis, which would be the correct derivative.

To implement this idea and prevent invalid product nodes from appearing, we modify the original DAG before it is differentiated by making the sum over the disappearing axis explicit. In nodes where the problem would occur, that is to say nodes where at least one index appears only in the index string of one of the input nodes, not in the output index string and where this input node contains the differentiation argument, we do the following. We identify the missing indices and append them to the output index string of the product node. This prevents an invalid product node from being created, but it also changes the node, since it now has more axes than before. To remedy this, we add another product node where we explicitly sum over these axes while multiplying with a vector of ones. This achieves the same result as the original DAG, but by making the sum over the axes explicit in another product node, we prevent the problematic case.

In our example, we would transform the original expression from  $A *_{(ij,j\to)} v$  to  $(A *_{(ij,j\to i)} v) *_{(i,i\to)} 1$ . Both the modified DAG and the derivative are visible in Figure 4.11.

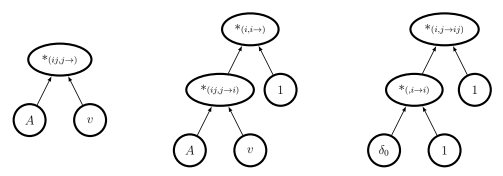


Figure 4.11: Left: DAG with a disappearing index i. Middle: Modified DAG where the sum over index i is made explicit. Right: Derivative of the middle DAG with respect to A.

## 5 Literature

## **Bibliography**

- [1] Sören Laue, Matthias Mitterreiter, and Joachim Giesen. "A Simple and Efficient Tensor Calculus". In: *The Thirty-Fourth AAAI Conference on Artificial Intelligence (AAAI-20)* (2020), pp. 4527–4534.
- [2] Sören Laue, Matthias Mitterreiter, and Joachim Giesen. "Computing Higher Order Derivatives of Matrix and Tensor Expressions". In: 32nd Conference on Neural Information Processing Systems (NeurIPS 2018) (2018).

# 6 Appendices

# 7 Declaration of autonomy