A multi-task learning-based optimization approach for finding diverse sets of material microstructures with desired properties and its application to texture optimization

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Abstract

The optimization along the chain processing-structure-properties-performance is one of the core objectives in data-driven materials science. In this sense, processes are supposed to manufacture workpieces with targeted material microstructures. These microstructures are defined by the material properties of interest and identifying them is a question of materials design. In the present paper, we address this issue and introduce a generic multi-task learning-based optimization approach. The approach enables the identification of sets of highly diverse microstructures for given desired properties and corresponding tolerances. Basically, the approach consists of an optimization algorithm that interacts with a machine learning model that combines multi-task learning with siamese neural networks. The resulting model (1) relates microstructures and properties, (2) estimates the likelihood of a microstructure of being producible, and (3) performs a distance preserving microstructure feature extraction in order to generate a lower dimensional latent feature space to enable efficient optimization. The proposed approach is applied on a crystallographic texture optimization problem for rolled steel sheets given desired properties.

Keywords— crystal plasticity, distance preserving feature extraction, machine learning, materials design, multi-task learning, multidimensional scaling, siamese neural networks, texture optimization
1 Introduction

1.1 Motivation

The demand for more and more specific and individually designed products with certain performance requirements has become a driving force in the world of manufacturing. For this reason, the optimization along the causal chain processing-structure-properties-performance \[1\] became a fast growing research topic in the field of integrated computational materials engineering (ICME) \[2\]. Nowadays, such optimization problems can be solved efficiently with the help of machine learning techniques \[3\]. On this background, in a previous work, we investigated the use of reinforcement learning for finding optimal processing routes in a simulated metal forming process aiming to produce microstructures with targeted crystallographic textures \[4\]. To bridge the remaining gap between microstructures and desired properties, we focus in this work on solving materials design problems. These are to identify appropriate material microstructures or microstructural features (e.g. the crystallographic texture) for given desired properties. It is thereby of particular importance to identify sets of near-optimal and preferably diverse microstructures in order to guarantee a robust design \[5\].

1.2 Paper structure

In the following we summarize the related work and point out the contribution of this paper. In Section \[2\] first, we describe the siamese multi-task learning and optimization approach. Then, we introduce the fundamentals in materials modeling that are needed for the purpose of this work. After that, in Section \[3\] the results are shown when applying the approach to a texture optimization problem for steel sheets. In Section \[4\] the presented results are discussed. Finally, in Section \[5\] we summarize our findings and give an outlook on further research.

1.3 Related work

A recent and very generic approach to solve materials design problems is the microstructure sensitive design (MSD) approach introduced in \[6\]. Following \[7\], MSD can be described by the seven steps. First, the properties of interest as well as candidate materials have to be defined. After that, a suitable microstructure definition is applied for these materials yielding a microstructure design space. On this basis, relevant homogenization relations are identified and applied over the whole design space. The resulting properties closure can be used to select desired properties, which are then mapped back to the microstructure design space in order to identify optimal microstructures. The last step of MSD aims to determine processes and processing routes needed to produce the identified microstructure.
The works by Adams et al. [6] and Kalidindi et al. [8] instantiate the MSD approach for texture optimization. The first one describes how optimal crystallographic textures can be identified in order to improve the deformation behavior of a compliant beam. In the latter, a similar approach is shown to optimize the crystallographic texture for the design of an orthotropic plate. The core of both approaches lies in the usage of a lower dimensional spectral representation of the orientation distribution, cf. [9]. For more complex microstructure representations, like two-point correlations, feature extraction methods can be applied to reduce the dimensionality. Methods that are for example used in the context of materials design are principal component analysis (PCA) [10, 11] and multidimensional scaling [12]. A general review of dimensionality reduction techniques can be found in [13].

Besides the MSD approach, also machine learning-based approaches for crystallographic texture optimization exist. [14] and [15] describe iterative sampling approaches that interact with crystal plasticity simulations aiming to identify crystallographic textures for given desired properties. Therefore, an initial set of texture-properties tuples is generated. Via supervised learning, significant features of the parameterized orientation distribution (and in [14] also regions) are identified that yield optimal or near-optimal solutions. Based on the identified features and regions, new texture-properties data points are sampled in order to get closer to the optima.

Another approach for identifying optimal textures is described in [16]. Therein, a real-coded genetic algorithm [17] is described that interacts with a crystal plasticity model in order to find optimal combinations of typical fcc rolling texture components (Cu, Brass, S, Cube and Goss) for given desired properties. The algorithm starts with an initial set of textures consisting of different fractions of these components. The set of textures evolves iteratively by combining them using operators such as mutation, crossover and selection [18].

Summarized, for the solution of microstructure design problems, a linkage from properties to microstructures is required. Such a linkage is often achieved by genetic or optimization algorithms that interact with numerical simulations. However, as these algorithms generally need a lot of function evaluations, it is not reasonable to apply them to complex numerical simulations directly. Instead, the performance can be increased by using numerically simpler surrogate models [19]. Typically, these are supervised learning models that learn the input-output relations of the numerical simulation under consideration.

To run optimization algorithms in combination with supervised learning models it is necessary to limit the region in which they operate to the region, which is covered by the training data. One way to achieve this is by training unsupervised learning methods on the input data, as it is for example done in [20] using support vector machines (SVM). From a machine learning perspective such an approach can be seen as anomaly detection. Anomaly
detection aims to separate data that is characteristically different from the
known data of the sample data set, which has been used for training. An
extensive overview of anomaly detection methods is given in [21]. Moreover,
[22] gives an overview on recent deep learning-based approaches for anomaly
detection, from which we want to point out neural network-based autoen-
coders [23], which fit especially well into multi-task learning (MTL) schemes
other than SVMs.

Autoencoder approaches assume that features of a data set can be mapped
into a lower dimensional latent feature space, in which the known data
points differ substantially from unknown data points. By backmapping into
the original space, anomalies can be identified by evaluating the reconstruc-
tion error, see for example [24]. In [24] it is also shown that autoencoder
networks are able to detect subtle anomalies, which cannot be detected by
linear methods like PCA. Furthermore, autoencoder networks require less
complex computations compared to a nonlinear kernel-based PCA.

1.4 Contribution

In the present paper, we introduce a generic MTL-based optimization ap-
proach to efficiently identify sets of microstructures, which are highly divers
and producible by a process. The approach is based on an optimization al-
gorithm interacting with a machine learning model that combines MTL [25]
with siamese neural networks [26]. In contrast to [14, 15] and also to [16], in
our approach a surrogate model is set up in order to replace the numerical
simulation, which maps microstructures to properties. The microstructure-
properties mapping can be executed efficiently by means of the surrogate
model within the optimization procedure.

To address the issue of a producibility, we include a neural network in the
MTL structure, which estimates the validity of a microstructure in the sense
of being producible. The efficiency of the optimization is further increased
by transforming the microstructure representation into a lower dimensional
latent feature space, which is formed by a non-linear data-driven autoen-
coder. The resulting lower dimensional latent feature space delivers the
input for the three neural networks: The first network maps the features to
properties (surrogate model), the second network estimates the validity and
the third network is the decoder-part of the autoencoder. As learning takes
place simultaneously for the encoder and the attached tasks, it is ensured
that the lower dimensional feature space is optimal for all tasks. In addition,
we enforce the latent feature space to preserve microstructure distances by
employing a siamese neural network and multidimensional scaling. On this
basis, we force the optimizer to find a diverse set of solutions in the latent
feature space.
2 Methods

2.1 Materials design via siamese multi-task learning (SMTL) and optimization

2.1.1 General Concept

First of all, we present the general concept of our MTL-based optimization approach. The approach can be applied to general materials design problems and starts by defining the desired properties and corresponding tolerances. This defines a target region, for which the approach is supposed to identify a diverse set of microstructures. The approach is schematically depicted in Fig. 1 and basically consists of three components: optimizer, microstructure-properties mapping (m-p-m) and validity-prediction (v-p). The optimizer generates candidate microstructures that minimize the combined costs, which result from evaluations based on the m-p-m and v-p components.

Figure 1: General concept of the MTL-based optimization approach to solve materials design problems.

The m-p-m component assigns properties to a candidate microstructure. The deviation of the assigned properties to the target region determines the cost. In general, the m-p-m component can be realized by a numerical simulation. However, since numerical simulations are computationally expensive, a surrogate model is used instead. The surrogate model is realized by a regression model that learns the relations from a priori generated microstructure-properties data.

The v-p component is realized by an anomaly detection method which
Figure 2: Schematic illustration of a set of generated microstructures in a structure space $s_1, s_2$. The valid region is the part of structure space, which is supported by the sampled data. 'Unknown’ microstructures are located outside the valid region.

determines the validity of a candidate microstructure by comparing it to the set of valid microstructures. The $v-p$ component returns a value that can be seen as an estimate of a candidate microstructure being an element of the microstructure set under consideration. This is for example the set, which can be produced by a dedicated process (e.g. rolling). The value returned by the $v-p$ component defines the validity cost and drives the optimizer solution to a valid microstructure region, which is illustrated in Fig. 2. Besides, such a microstructure region can also be identified by a further optimization algorithm that interacts with a numerical simulation of the dedicated process, however, such an approach suffers from high computational costs.

The two components $m-p-m$ and $v-p$ can be realized by training two separate machine learning models. However, when the training procedures are isolated from each other, the models are not able to mutually access information already learned by the other model. Therefore, we combine the two components as tasks into one MTL model [25]. Both tasks have a common backbone (the feature extraction part of a network) and different heads (feature processing part of a network) operating on the backbone output. The backbone output vectors form the so-called latent feature space. The proposed MTL approach furthermore uses the backbone as an encoder network of an autoencoder, where the decoder is also attached to the latent feature space with the purpose to reconstruct the input pattern of the backbone. This is achieved by adding the reconstruction of the microstructures from
the latent feature space as a third task. In the MTL approach, all three tasks are represented by a single neural network-based model. The weights of the model are trained simultaneously based on a combined loss function. After training the MTL model, the optimizer can operate very efficiently in the lower dimensional latent feature space. The remainder of this section presents the optimization approach and the MTL approach in detail, as well as an extension based on siamese neural networks [26] to enforce the representation of microstructures in the latent feature space to preserve the microstructure distances in the original representation space.

2.1.2 Multi-task Learning (MTL)

The MTL processing scheme (shown in Fig. 3) starts with an encoder network which extracts significant features by mapping the microstructure space \( x \in \mathbb{R}^K \) into a lower dimensional latent feature space \( z \in \mathbb{R}^M \) via the learned function

\[
    z = f_{\text{enc}}(x, \theta_{\text{enc}}),
\]

in which the encoder network is parameterized by its weight values \( \theta_{\text{enc}} \).

All three previously described tasks are attached to the encoder in the form of feedforward neural networks. Besides, the encoder can be easily adapted to higher dimensional microstructure representing data types like images (EBSD or micrograph images) or three dimensional microstructure data by using for example convolutional neural networks (see [27]), which are used for example in [28] in the materials sciences domain. To train the MTL model, a loss function that combines all the three tasks is needed. This is achieved by a function that cumulates the loss terms of the three tasks \( \mathcal{L}_{\text{regr}} \) (regression loss), \( \mathcal{L}_{\text{recon}} \) (reconstruction loss) and \( \mathcal{L}_{\text{valid}} \) (validity loss), and weights them using \( \mathcal{W}_{\text{regr}} \), \( \mathcal{W}_{\text{recon}} \) and \( \mathcal{W}_{\text{valid}} \) to allow for prioritization. The total loss function is defined as

\[
    \mathcal{L}_{\text{MTL}} = \mathcal{W}_{\text{regr}} \mathcal{L}_{\text{regr}} + \mathcal{W}_{\text{recon}} \mathcal{L}_{\text{recon}} + \mathcal{W}_{\text{valid}} \mathcal{L}_{\text{valid}} + \lambda R(\theta),
\]

where \( R(\theta) \) is a regularization term that is used to prevent overfitting with the hyperparameter \( \lambda \) defining the strength of the regularization (also known as weight decay, see [29] and [30]). Each of the feedforward neural networks is parameterized by the respective weight values \( \theta_{\text{enc}}, \theta_{\text{regr}}, \theta_{\text{recon}} \) and \( \theta_{\text{valid}} \), which are adjusted simultaneously during training and altogether form the weight vector \( \theta \). In the following we will introduce the three individual loss terms.

1. The forward mapping of the latent feature vector \( z \) to the properties vector \( \hat{p} \in \mathbb{R}^N \) is represented by the learned function

\[
    \hat{p} = f_{\text{regr}}(z, \theta_{\text{regr}}) = f_{\text{regr}}(f_{\text{enc}}(x, \theta_{\text{enc}}), \theta_{\text{regr}}).
\]

7
2. The decoder network, which is responsible for the reconstruction, transforms the latent feature vectors $z$ back to the original microstructure space:

$$x' = f_{\text{recon}}(z, \theta_{\text{recon}}) = f_{\text{recon}}(f_{\text{enc}}(x, \theta_{\text{enc}}), \theta_{\text{recon}}).$$

The reconstruction loss is defined on the basis of a distance measure between two microstructural feature vectors $\text{dist}(x, x')$:

$$L_{\text{recon}}(x, x') = \text{dist}(x, x').$$

The distance measure between depends on the microstructure representation and has to be chosen appropriately.

3. On the basis of the latent feature space, an extra autoencoder network is set up transforming $z \in \mathbb{R}^M$ into an even lower-dimensional feature
sub-space \( s \in \mathbb{R}^S \) with \( S < M \) and transforming back to \( z' \in \mathbb{R}^M \) via

\[
z' = f_{\text{valid}}(z, \theta_{\text{valid}}) = f_{\text{valid}}(f_{\text{enc}}(x, \theta_{\text{enc}}), \theta_{\text{valid}}).
\]

The validity loss is defined by the mean squared error between \( z \) and \( z' \):

\[
\mathcal{L}_{\text{valid}}(z, z') = \frac{1}{M} \sum_{i=1}^{M} (z_i - z'_i)^2.
\]

### 2.1.3 Distance preserving feature extraction using siamese neural networks

The above described MTL approach is used in combination with an optimizer that searches for candidate microstructures with desired properties in the latent feature space. However, our approach aims to identify a diverse set of microstructures with high diversity. For the diversity quantification a distance measure in the latent feature space is required. The MTL approach as defined above, is not able to preserve the distances of the original space in the latent feature space. In order to construct a distance preserving latent feature space, the MTL is embedded in a siamese neural network \[26, 31\], which we will describe next.

Siamese neural networks consist of two identical networks, which share weights in the encoder part, see Fig. 4. Both networks embed different microstructures \( \vec{x}_L \) and \( \vec{x}_R \) as \( \vec{z}_L \) and \( \vec{z}_R \) in the latent feature space which is finally processed by two identical MTL networks. The distance preservation is enforced by defining a distance preservation loss \( \mathcal{L}_{\text{pres}} \) that minimizes the difference between the distance of two different input microstructures in the original space \( \text{dist}(x_L, x_R) \) and the corresponding distance in the latent feature space \( \text{dist}(z_L, z_R) \), with \( x_L \neq x_R \)[32]:

\[
\mathcal{L}_{\text{pres}} = (\text{dist}(x_L, x_R) - \text{dist}(z_L, z_R))^2,
\]

while \( \text{dist}(x_L, x_R) \) and \( \text{dist}(z_L, z_R) \) are not necessarily the same distance measures. Applying such loss terms leads to multi dimensional scaling, see [33] and [34]. Using the distance preservation loss \( \mathcal{L}_{\text{pres}} \), the MTL loss function, defined in Eq. 2 is extended by the weighted preservation loss \( \mathcal{W}_{\text{pres}} \mathcal{L}_{\text{pres}} \) to

\[
\mathcal{L}_{\text{SMTL}} = \mathcal{W}_{\text{regr}} \mathcal{L}_{\text{regr}} + \mathcal{W}_{\text{recon}} \mathcal{L}_{\text{recon}}
+ \mathcal{W}_{\text{valid}} \mathcal{L}_{\text{valid}} + \mathcal{W}_{\text{pres}} \mathcal{L}_{\text{pres}}
+ \lambda R(\theta).
\]

The SMTL approach delivers a function which can map a microstructure representation in the latent feature space on properties. Now an optimizer can operate on a low dimensional feature space to find microstructures with desired properties. The SMTL framework also allows to reconstruct the original representation of microstructures, to assess the distances between them and to validate them in the latent feature space.
2.1.4 Microstructure optimizer

The microstructure optimization with respect to desired properties uses the distance preserving SMTL framework with the tasks microstructure-property-mapping, validity-prediction and reconstruction. The optimization minimizes a loss function, which consists of the cost terms $C_{\text{prop}}$, $C_{\text{valid}}$ and $C_{\text{divers}}$ and the corresponding weights $V_{\text{prop}}$, $V_{\text{valid}}$ and $V_{\text{divers}}$:

$$F = V_{\text{prop}} C_{\text{prop}} + V_{\text{valid}} C_{\text{valid}} + V_{\text{divers}} (1 + C_{\text{divers}}).$$ (11)

$C_{\text{prop}}$, $C_{\text{valid}}$ and $C_{\text{divers}}$ denote the property, validity and diversity cost terms, respectively. While the property cost term drives the candidate microstructures to lie inside a specified target properties region, the validity cost aims that the optimizer operates inside the region of valid microstructures and the diversity cost ensures that candidate microstructures differ.
from each other. To minimize the loss function we use genetic algorithms, which generate a population set of \( P \) candidate microstructures \( \tilde{z}^* \) in the latent feature space in every iteration. The three cost terms are described in more detail in the following.

1. The property cost is defined by the mean squared error between the desired properties and the predicted properties from the SMTL regression model:

\[
C_{\text{prop}} = \frac{1}{N} \sum_{i=1}^{N} (\tilde{C}_{\text{prop},i})^2.
\]  

(12)

If one of the predicted properties lies inside the target region, the cost \( \tilde{C}_{\text{prop},i} \) equals 0. Otherwise, \( \tilde{C}_{\text{prop},i} \) equals the minimum squared distance from the predicted properties to the target region borders.

2. The validity prediction is used to assess whether an identified candidate microstructure is likely to be represented by the sample data set. The validity cost is defined by

\[
C_{\text{valid}} = \max(A - \xi_{\text{valid}}, 0),
\]  

(13)

in which \( \xi_{\text{valid}} \) is a threshold to define the maximum tolerated reconstruction error for valid textures and \( A \) denotes the anomaly score

\[
A = \frac{1}{M} \sum_{i=1}^{M} (z_i^{*} - z_i^{**})^2.
\]  

(14)

3. The diversity cost is based on the sum of the distances between the candidate microstructure \( z^* \) in the latent feature space and every other microstructure in the population:

\[
C_{\text{divers}} = - \sum_{i=1}^{P} \text{dist}(z_i^{*}, z^*),
\]  

(15)

in which for \( \text{dist}(z_i^{*}, z^*) \) the same distance measure has to be used as for the latent feature vectors in Eq. 9.

2.2 Materials science fundamentals

2.2.1 Representation of crystallographic texture

Crystallographic texture is typically described by the orientation distribution function, which is defined by

\[
f(g)dg = \frac{V(g)}{V},
\]  

(16)
for an orientation \( g \) (a point in \( SO(3) \)) and the volume \( V(g) \) in \( SO(3) \). The orientation distribution function \( f(g) \) often underlies specific symmetry conditions, for which various regions in \( SO(3) \) are equivalent. Therefore, depending on the symmetries, orientations can be mapped into an elementary region of \( SO(3) \), the so-called fundamental zone. The orientation distribution function on the basis of the orientations mapped into the fundamental zone is then indistinguishable from the original orientation distribution function. Rolling textures, for example, underlie a cubic crystal and an orthorhombic sample symmetry, for which 96 elementary regions exist \[35\].

A popular way to represent the orientation distribution function is by approximating it via generalized spherical harmonic functions \[9\]. Yet, as there is no straightforward way to measure the distance between two orientation distribution functions in terms of generalized spherical harmonics, we make use of the orientation histogram-based texture descriptor, which is introduced in \[4\]. Therefore, the cubic fundamental zone is discretized into a set \( O \) of \( j \) nearly uniform distributed orientations \( o_j \). For each individual orientation \( g \) in a set of orientations \( G \), a weight vector \( w_g \) is constructed via a soft-assignment approach

\[
 w_g = \begin{cases} 
 \frac{\Phi(g, o_j)}{\sum_{o_i \in N_l} \Phi(g, o_i)}, & \text{if } o_j \in N_l \\
 0, & \text{else}
\end{cases}
\] (17)

where \( N_l \) is the set of \( l \) nearest neighbor orientations of \( g \) in terms of the orientation distance \( \Phi \). The orientation distance between two orientations \( g \) and \( o \) is defined by

\[
 \Phi = \min \Phi(\overline{g}, \overline{o}).
\] (18)

where \( \overline{g} \) and \( \overline{o} \) is from the set of all equivalent orientations of \( g \) and \( o \) in terms of cubic crystal symmetry. The orientation distance measure in \( SO(3) \) is defined as

\[
 \Phi(q_g, q_o) = \min(||q_g - q_o||, ||q_g + q_o||),
\] (19)

where \( q_g \) and \( q_o \) are the quaternion representations of the orientations \( g \) and \( o \) \[36\].

On this basis, the weight vector for the orientation histogram \( b \) can be calculated by a volume average of the weight vectors of the individual orientations

\[
 b = \frac{1}{V} \sum_j V(o_j)w_{o_j}.
\] (20)

The distance between two orientation distribution functions can then be measured via any kind of histogram-based distance measure, such as the Chi-Squared distance \[37\]

\[
 \chi^2(b_1, b_2) = \sum \frac{(b_1 - b_2)^2}{b_1 + b_2}.
\] (21)
Table 1: Definition of the fibers of bcc rolling textures following [41]

| Fiber | Location |
|-------|----------|
| α     | from \{001\}<110> to \{111\}<110>, parallel to RD |
| γ     | from \{111\}<110> to \{111\}<112>, parallel to ND |
| η     | from \{001\}<100> to \{011\}<100>, parallel to RD |
| ϵ     | from \{001\}<110> to \{111\}<112>, parallel to TD |
| β     | from \{112\}<110> to \{11\}4 \{11\}8 <4 4 \{11\}> |

The set of nearly uniform distributed orientations \( O \), needed for the histogram-based texture descriptor, can be generated using the algorithm described in [38], which is implemented in the software neper [39]. For the purpose of this study, we sample 512 nearly uniform distributed orientations over the cubic fundamental zone and chose a soft assignment of \( l = 3 \).

### 2.2.2 Crystallographic texture of steel sheets

After rolling body centered cubic (bcc) materials, typically so-called fiber textures are formed. Following [40], these textures are composed of the five fibers \( \alpha, \gamma, \eta, \epsilon, \) and \( \beta \), which are defined in detail in Tab. 1. Among these fibers, the \( \alpha \) and \( \gamma \) fiber are most prominent [41], whereas the presence of the \( \beta \) fiber is only reported from theoretical predictions [42]. In order to generate a data base of (artificial) rolling textures, in this work, a 25-parameter model is used, as it is proposed in [43] to describe steel sheet textures. The model is based on textures that are composed of the fibers \( \alpha, \gamma, \) and \( \eta \).

As the \( \eta \)-fiber is not always present in steel sheet textures, we limit ourselves to textures that consist of an \( \alpha \) and \( \gamma \) fiber. Therefore, 6 of the 25 parameters can be neglected. The texture model describes the orientation distribution function as a set of weighted Gaussian distributions placed along the fibers. The model parameters \( D_i \) are listed in Tab. 2 and define the standard deviations and the mean values of the distributions based on the fiber thickness and the shifts from their ideal positions. Furthermore, the model parameters define the weights of the distributions among each other based on the fiber intensity.

To construct the set of Gaussian distributions, the seven base distributions from Tab. 2 are placed at their ideal positions with respect to the shifts. Between these seven distributions, further distributions are placed with a distance of about \( 3^\circ \) to each other, leading to overall 41 Gaussians. Their weights \( w_i \) and the values for the standard deviation \( \sigma_i \) and mean value \( \mu_i \) are interpolated linearly based on the values of the two neighboring base distributions. This yields a set of Gaussian distributions.
Table 2: Definition of the parameters $D_i$ of the texture model, cf. [43]. The ideal position is given in Bunge euler angles $[^\circ]$.

| Ideal pos. | Intens. | Std $\varphi_2$ | Std $\varphi_1$ | Shift $\varphi_1$ | Shift $\varphi_2$ |
|------------|---------|-----------------|-----------------|-------------------|------------------|
| a1 0, 0, 45 | $D_1$   | $D_7$           | $D_7$           | 0                 | 0                |
| a2 0, 30, 45| $D_2$   | $D_8$           | $D_{13}$        | 0                 | $D_{15}$        |
| a3 0, 55, 45| $D_3$   | $D_9$           | $D_9$           | 0                 | $D_{16}$        |
| a4 0, 70, 45| $D_4$   | $D_{10}$        | $D_{10}$        | 0                 | $D_{17}$        |
| a5 0, 90, 45| $D_5$   | $D_{11}$        | $D_{11}$        | 0                 | $D_{18}$        |
| a6 15, 55, 45| $D_6$  | $D_{12}$        | $D_{12}$        | 0                 | $D_{19}$        |
| a7 30, 55, 45| $D_7$  |                |                |                   |                  |

$N_i(\mu_1, \sigma_1), ..., N_{41}(\mu_{41}, \sigma_{41})$. The orientation distribution function $f(g)$ is defined by the normalized sum of this set:

$$f(g) = \frac{1}{\sum_i w_i \sum_{i=1}^n w_i N_i(\mu_i, \sigma_i)}.$$  (22)

Based on this definition, discrete orientations can be sampled. In the following, we denote the set of orientations as $G$. As $f(g)$ is defined in the cubic-orthorhombic fundamental zone, it is necessary to add the equivalent orientations regarding the orthorhombic sample symmetry to the set of discrete orientations. This is done by applying rotation operations $g_s$ on each orientation $g_i$ in $G$

$$g_i^{\text{equiv}} = g_s g_i.$$  (23)

The rotation operations $g_s$ for orthorhombic sample symmetry can be found in [35].

2.2.3 Material model

The sheet metal properties which we focus on in this study are the Young’s moduli and the R-values at 0, 45 and 90 degree to rolling direction. In this study, the properties are calculated by applying uniaxial tension on a crystal plasticity-based material model. As time efficiency is essential for the generation of data, a material model of Taylor-type is implemented, as it is described in [4].

The Taylor-type material model is based on the volume averaged stress of a set of $n$ crystals [44]:

$$\overline{T} = \frac{1}{V} \sum_{i=1}^n T^{(i)} V^{(i)}.$$  (24)
In the above equation, \( \mathbf{T} \) denotes the Cauchy stress tensor, which can be derived by the stress tensor in the intermediate configuration, given by

\[
\mathbf{T}^* = \frac{1}{2} \mathbf{C} : (\mathbf{F}_e^T \cdot \mathbf{F}_e - \mathbf{I}),
\]

with the second order identity tensor \( \mathbf{I} \) and the fourth order elastic stiffness tensor \( \mathbf{C} \). The elastic constants \( C_{11}, C_{12} \) and \( C_{44} \) are set to 218.37, 131.13 and 105.34 GPa, respectively \[45\]. \( \mathbf{F}_e \) is the elastic part of the deformation gradient \( \mathbf{F} \) and can be calculated by a multiplicative decomposition

\[
\mathbf{F} = \mathbf{F}_e \cdot \mathbf{F}_p.
\]

The intermediate stress tensor can be converted into Cauchy stress using the relation

\[
\mathbf{T}^* = \mathbf{F}_e^{-1} \cdot (\det(\mathbf{F}_e) \mathbf{T}) \cdot (\mathbf{F}_e)^{-T}.
\]

To describe the evolution of the plastic deformation, the plastic part of the velocity gradient \( \mathbf{L}_p \) is considered by

\[
\mathbf{L}_p = \dot{\mathbf{F}}_p \cdot \mathbf{F}_p^{-1},
\]

and the flow rule \[46\]

\[
\mathbf{L}_p = \sum_n \dot{\gamma}^{(n)} \mathbf{m}^{(n)} \otimes \mathbf{n}^{(n)},
\]

where \( \dot{\gamma}^{(n)} \) denotes the shear rates on the active slip systems \( \eta \), defined by the slip plane normal \( \mathbf{n}^{(n)} \) and the slip direction \( \mathbf{m}^{(n)} \). For bcc materials, the slip system families in terms of Miller index are \{110\}<111>, \{112\}<111>, and \{123\}<111>, while the latter is neglected due to simplicity.

The shear rates are defined by a phenomenological power-law \[47\]:

\[
\dot{\gamma}^{(n)} = \dot{\gamma}_0 \left| \frac{\tau^{(n)}}{\tau^{(n)}} \right|^{1/m} \text{sign}(\tau^{(n)}),
\]

where \( \tau^{(n)} \) is the slip system resistance, \( \dot{\gamma}_0 \) the reference shear rate and \( m \) the shear rate sensitivity. Here, \( \dot{\gamma}_0 \) and \( m \) are set to 0.001 sec\(^{-1}\) and 0.0125, respectively, \[48\]. Following Schmid’s law, the resolved shear stress on slip system \( \tau^{(n)} \) is given by

\[
\tau^{(n)} = ((\mathbf{F}_e^T \cdot \mathbf{F}_e) \cdot \mathbf{T}^*) : (\mathbf{m}^{(n)} \otimes \mathbf{n}^{(n)}),
\]

and the evolution of the slip system resistance is defined by

\[
\dot{\tau}^{(n)} = \frac{d\tau^{(n)}}{d\Gamma} \sum_{\xi} q_{\xi} |\dot{\gamma}^{(\xi)}|. \]
The matrix $q_{\gamma\xi}$ describes the ratio between self and latent hardening. It consists of diagonal elements equal to 1.0 and off-diagonal elements $q_1$ and $q_2$, cf. [19]. Both, $q_1$ and $q_2$, are set to 1.4 [17]. Further, we need to model the hardening behavior, which is realized by an extended Voce-type model [50]:

$$\hat{\tau}(\eta) = \tau_0 + (\tau_1 + \vartheta_1 \Gamma)(1 - e^{-\Gamma \vartheta_0 / \tau_1}).$$  \hspace{1cm} (33)

The material dependent parameters are calibrated to DC04 steel\textsuperscript{1} and are $\tau_0 = 94.9$ MPa, $\tau_1 = 50$ MPa, $\vartheta_0 = 258$ MPa and $\vartheta_1 = 32.8$ MPa [51]. The accumulated plastic shear is defined by

$$\Gamma = \int_0^t \sum_\eta |\hat{\tau}(\eta)| \, dt. \hspace{1cm} (34)$$

3 Results

3.1 Texture-property data set

For training, 50000 sets of 2000 discrete orientations are sampled via Latin Hypercube Design [52], based on Eq. 22. In order to have an independent test set, further 10000 sets are generated randomly. The ranges inside which the parameters of the texture model vary are defined such that typical bcc rolling textures found in literature can be represented, cf. [53, 54, 55, 41, 56, 48, 57]. The parameter ranges are listed in Tab. 3. In addition, to evaluate the anomaly detection, a set of artificial textures is needed, which slightly differ from the generated rolling textures. For this purpose, 10000 anomalies are generated by shifting the $\alpha$-fiber (i.e. the ideal position of $a_1$, $a_2$, $a_4$ and $a_5$) about 20 degrees in $\varphi_1$-direction.

Moreover, we want to validate the texture-property-mapping and the validity-prediction on experimental data. For this purpose, an experimentally measured texture of cold rolled DC04 steel from [58] is used. Based on this measurement, an orientation distribution function is approximated via the MATLAB toolbox mtex [59], rotated into its symmetry axis assuming orthorhombic sample symmetry and mirrored. To visualize the $\alpha$- and $\gamma$-fiber of the orientation distribution, an intersection plot of the euler space at $\varphi_2 = 45^\circ$ is depicted in Fig. 5.

3.2 Validation of SMTL

In this study, the individual tasks of the SMTL model are realized via feed-forward neural networks with $tanh$ activation functions to obtain features between $-1$ and $+1$ in the latent feature space. The SMTL model is implemented based on the Python TensorFlow API [60]. The base network of

\textsuperscript{1}experiments performed at IUL Dortmund during DFG project Graduate School 1483 [51]
Table 3: Parameter ranges for $D_i$

| Intensity | $D_1$ | $D_2$ | $D_3$ | $D_4$ | $D_5$ | $D_6$ |
|-----------|-------|-------|-------|-------|-------|-------|
| min       | 1/6   | 1/6   | 1/6   | 0     | 1/6   | 1/6   |
| max       | 1/3   | 2/3   | 2/3   | 1/3   | 1     | 1     |
| Std       | $D_7$ | $D_8$ | $D_9$ | $D_{10}$ | $D_{11}$ | $D_{12}$ | $D_{13}$ |
| min       | 5/3   | 5/3   | 5/3   | 5/3   | 5/3   | 5/3   | 5/3   |
| max       | 45/3  | 35/3  | 35/3  | 30/3  | 30/3  | 30/3  | 35/3  |
| Shift     | $D_{14}$ | $D_{15}$ | $D_{16}$ | $D_{17}$ | $D_{18}$ | $D_{19}$ |
| min       | −5    | −10   | −10   | −5    | −10   | −10   |
| max       | 10    | 10    | 5     | 10    | 10    | 10    |

Figure 5: $\varphi_2 = 45^\circ$ section of the orientation distribution function to visualize the $\gamma$-fiber of the colled rolled DC04 steel texture.
Table 4: Used hyperparameters

| Hyperparameter   | Value |
|------------------|-------|
| Optimizer        | Adam  |
| Learning rate    | 0.001 |
| Weight decay     | 1e-6  |
| Batch size       | 256   |

The siamese architecture is illustrated in Fig. 6. The Glorot Normal method [61] is used for weight initialization. In order to adjust the hyperparameters, a random search method [62] is applied using 5-fold cross-validation. The best model configuration that was found is shown in Tab. 4. We use the Chi-Squared distance introduced in Eq. 21 as the distance measure in the input space. In the latent latent feature space we use the sum of squared errors (SSE) between two vectors $z_1$ and $z_2$ as distance measure

$$
SSE(z_1, z_2) = \sum_{i=1}^{M} (z_{1,i} - z_{2,i})^2.
$$

The SMTL model is trained for 200 epochs, while the best intermediate result of the test set is retained, which can be interpreted as a form of early stopping [63]. Before the model training is executed, the loss terms are scaled to values between 0 and 1 in order to make them comparable. The following weights for the scaled loss terms were found to be appropriate from hyper parameter optimization: $\mathcal{W}_{\text{regr}} = 0.05$, $\mathcal{W}_{\text{recon}} = 0.05$, $\mathcal{W}_{\text{valid}} = 0.05$ and $\mathcal{W}_{\text{pres}} = 0.85$.

The results for the texture-property-mapping and the distance preservation are shown in Tab. 5, in which the regression errors $\text{MAE}_E$ and $\text{MAE}_r$
Table 5: Results for varying numbers of latent features (LF) of the texture-property-mapping and the distance preservation applied to the artificially generated textures and experimentally measured texture. Regr. (ession error) MAE_E is given in \([\text{MPa}]\), Regr. (ession error) MAE_r in [-] and Pres. (erve quality) \(R^2\) in [%].

| LF | Artificial textures | Experimental texture |
|----|---------------------|----------------------|
|    | Regr. MAE_E | Regr. MAE_r | Pres. \(R^2\) | Regr. MAE_E | Regr. MAE_r |
| 20 | 153     | 0.03       | 98.2   | 596     | 0.04       |
| 18 | 183     | 0.03       | 98.0   | 773     | 0.11       |
| 16 | 162     | 0.03       | 97.8   | 907     | 0.05       |
| 14 | 193     | 0.03       | 97.0   | 1282    | 0.07       |
| 12 | 215     | 0.04       | 95.4   | 1468    | 0.13       |
| 10 | 238     | 0.05       | 92.2   | 1463    | 0.13       |
| 8  | 335     | 0.06       | 85.7   | 1575    | 0.17       |
| 6  | 390     | 0.07       | 72.4   | 1554    | 0.15       |
| 4  | 664     | 0.10       | 34.2   | 2768    | 0.12       |

denote the mean absolute error between the true and predicted Young’s moduli and R-values depending on the dimension of the latent feature space \(z\). The quality of the distance preservation is measured by the coefficient of determination \(R^2\), between the distances of two input textures and their corresponding latent feature vectors \(R^2(\chi^2(x_L, x_R), SSE(z_L, z_R))\). It is shown that texture-property-mappings with an adequate prediction quality can be achieved by extensively reducing the dimensionality of the latent feature space. However, regarding the distance preservation quality, a lower bound of at least 10 latent features can be identified, below which the distance preservation is unsatisfactory. Additionally, the texture-property-mapping is evaluated on the experimentally measured texture and the corresponding properties. The results are listed in Tab. 5. It can be seen that a satisfactory prediction quality (Regr. MAE_E \(\leq\) 1000 MPa and Regr. MAE_r \(\leq\) 0.1) can only be achieved for at least 16 latent features.

On the basis of this 16-dimensional feature space, the validity-prediction is then evaluated. The anomaly scores for the textures in the test set and for the artificially generated anomalies are shown in Fig. 7. It can be seen, that the anomalies can be separated in a sufficient manner from the textures in the test set.

### 3.3 Rolling texture identification

To validate the texture identification, we define two target regions in the property space, see Fig. 8. The first one is defined by the properties of the experimentally measured texture, which lies in a sparsely populated re-
Figure 7: Histograms of the anomaly scores for the data from the test set and the set of artificially generated anomalies. The anomaly scores are based on the model that uses 16 latent features.

Table 6: Center points of the two target regions (TR). The Young’s moduli $E$ are given in MPa, the R-values $r$ in [-].

| TR | $E_{00}$ | $E_{45}$ | $E_{90}$ | $r_{00}$ | $r_{45}$ | $r_{90}$ |
|----|---------|---------|---------|---------|---------|---------|
| 1  | 223145  | 207148  | 216599  | 1.01    | 1.7     | 0.4     |
| 2  | 223000  | 213000  | 222000  | 1.3     | 2.2     | 2.2     |

gion and is labeled as Target Region 1. As a consequence of its location in the sparsely populated region, the anomaly score of this texture is 0.0099 and in the transition zone shifted towards the generated anomalies (cf. Fig. 7). It is of interest if the optimizer is generally able to find a whole set of microstructures with properties in this region. The second target region represents a densely populated region located near the center of the properties point cloud and is labeled as Target Region 2. The center of the each target region is listed in Tab. 6. The target regions are defined by adding a tolerance of ±1000 MPa to the Young’s moduli and ±0.10 to the R-values. As a baseline, we collect all data points from the training set, that lie inside the target regions. In Target Region 1 only two textures can be found, whereas in Target Region 2 13 textures can be found.

To identify a diverse set of textures, we use the optimization algorithm JADE [65], which is an extension of the Differential Evolution algorithm [66]. Before starting the optimization via JADE, an initial population has to be selected, where 100 textures are sampled from the test set, which are
Figure 8: Density of the training set projected on different planes of the property space: the Young’s modulus at (a) 0 vs. 90 degree and (b) 45 vs. 90 degree for the R-values at (c) 0 vs. 90 degree and (d) 45 vs. 90 to rolling direction. The orange and red squares mark the projections of Target Region 1 and Target Region 2, respectively. The green dots show the projected samples from the training set that lie inside the target region.
approximately uniformly distributed over the property space. For the cost function, defined in Eq. 11, we use the weights $\mathcal{V}_{\text{prop}} = 0.90$, $\mathcal{V}_{\text{valid}} = 0.03$ and $\mathcal{V}_{\text{divers}} = 0.07$ and scale $C_{\text{props}}$ and $C_{\text{divers}}$ to values between 0 and 1 based on the selected 100 initial textures. The threshold $\xi_{\text{valid}}$ is set to 0.01 based on the maximum anomaly score in the data set, cf. Fig. 7. The optimization is performed for 300 iterations with a fixed population size of 100. During the optimization, all valid textures that fulfill the target properties are collected, according to the texture-property-mapping. Based on the results from the previous section, we use the trained SMTL-model with a 16-dimensional latent feature space. The resulting textures for each target region are discussed in the following.

Target region 1

Our approach is able to find a diverse set of textures that meet the property requirements of Target Region 1, according to the texture-property-mapping. Fig. 9 depicts the mutual distances in the latent feature space between all the found textures and between the two baseline textures. It is shown, that the set of identified textures contains 1315 diverse textures in contrary to only two in the baseline set. In order to compare the results to the experimentally measured texture, the closest texture to the center point of Target Region 1 is depicted in Fig 10 as a section through euler space at $\varphi_2 = 45^\circ$. By comparing the two textures, it can be seen that they are roughly the same in terms of the magnitude of the intensities and the shape of the $\alpha$- and $\gamma$-fibers. However, they also show differences in terms of smoothness and the location of the intensity peaks.

Target region 2

Compared to Target Region 1, an even more diverse set of 221 textures can be identified for Target Region 2, which can be seen in the histogram of the mutual distances in Fig. 11. To get an idea of the differences between the textures, two exemplary textures are plotted in Fig. 12 as a section through the ODF in the euler space at $\varphi_2 = 45^\circ$. It can be seen that the $\alpha$- and $\gamma$-fiber of both textures differ significantly in terms of intensity. However, the locations of the intensity peaks and the thickness of the $\alpha$- and $\gamma$-fiber are similar.

4 Discussion

The results presented in Section 3.2 show that the two tasks texture-property-mapping and validity-prediction are solved by the SMTL model. To achieve a sufficient prediction quality for both tasks in the test set as well as for the experimentally measured texture, a minimum dimensionality of the latent
Figure 9: Histogram of pairwise SSE distances of the set of identified textures and the baseline set for Target Region 1. The distance between the two textures from the baseline set is indicated by the dashed line.

Figure 10: Texture that yields properties which are closest to the center of Target Region 1. The plot shows the $\varphi_2 = 45^\circ$ section of the orientation distribution function.
Figure 11: Histogram of mutual distances of the set of identified textures and the baseline set for *Target Region 2*.

Figure 12: Two exemplary textures from the set of identified textures. Both plots show $\varphi_2 = 45^\circ$ sections of the respective orientation distribution functions.
feature space is needed, where also the dimensionality requirements of the siamese distance preservation goal has to be considered. 16 latent features were found to be sufficient for our example task of cold rolled steel. However, the prediction error for the experimentally measured texture is higher than for the test set using the same latent feature space dimensionality. This may be explained by the fact that the corresponding property is in a texture space region with low sampling density and the model therefore is not well supported by data. This results also in an instability of the model quality depending on the dimensionality of the latent feature space in this region. This instability can be seen by studying the R-value in Tab. \[\text{Tab. 5}\]. By choosing the latent feature space size of 16, also the results for the experimentally measured texture are satisfactory, especially keeping in mind that the experimentally measured texture differs naturally from the simulated data and additionally lies in a sparsely sampled region, cf. Target Region 1 in Fig. \[\text{Fig. 8}\].

Due to the sparsity of Target Region 1, the identification of textures in this region is challenging. Nevertheless, the optimization approach is able to identify a set of textures that contains more diverse individuals compared to the two baseline textures from the training set. Regarding the identified texture, which is closest to the experimentally measured texture in terms of properties, one can see that they are also similar in terms of crystallographic texture, what basically proofs the concept of our approach.

The most obvious difference between both textures is smoothness. The irregular distribution of intensity peaks of the identified texture is due to the resolution of the histogram-based texture descriptor. Also the orthorhombic sample symmetry is not represented locally. However, by increasing the resolution, these two issues can be solved. Furthermore, a higher resolution of the descriptor decreases the descriptor error, which reflects the deviation between the properties of the original texture and the properties of the texture described by the descriptor. However, the choice of resolution is a trade-off between accuracy and descriptor complexity, which is why we chose an intermediate resolution to proof our approach. Generally, with the use of the SMTL model and the incorporated feature extraction, the resolution is limited only by computational power.

Compared to Target Region 1, the identification task for Target Region 2 seems to be less challenging as the target region is located in a densely sampled region. However, as there already exists a proper set of diverse textures in the baseline, the main challenge is to outperform the baseline set in terms of diversity. Fig. \[\text{Fig. 11}\] shows that this general materials design problem (the identification of multiple equivalent microstructures) is accomplished by the optimization approach. This is exemplarily shown when comparing two of the identified textures in Fig. \[\text{Fig. 12}\] with each other: similar properties can be reached by quite different microstructures. The identification of such a highly diverse set of microstructures with similar properties is an important precondition to construct robust optimizing process control al-
gorithms, which can then choose among multiple optimal paths leading to desired properties.

5 Summary and outlook

In this work we present an approach to solve materials design problems. The approach is based on an optimization strategy that incorporates machine learning models for mapping microstructures to properties and for assessing the validity of input microstructures in the sense of the likeliness with the underlying data. To model these tasks, we use a siamese multi-task learning neural network (SMTL). Furthermore, we incorporate feature extraction in order to transform input microstructures to a lower dimensional latent feature space where the optimizer looking for microstructures with dedicated properties can efficiently operate.

By training the SMTL network with a dedicated loss function term, we are able to preserve the distances between microstructures in the original input space also in the latent feature space. The distance preservation allows to directly assess the diversity of the solution set found by the optimizer directly in the latent feature space and therefore enables optimizers to efficiently identify sets of diverse microstructures. By applying the approach to crystallographic texture optimization, we show the ability to identify diverse sets of textures that lie within given bounds of properties. Such sets of textures form the input of optimal processing control approaches like in [4].

In the present work, we applied our approach on data from mean-field simulations. The next step is to apply the approach on spatially resolved data from full-field simulations. The proposed methods can be easily extended for this task by modifying the encoder part of the SMTL network. However, the problem arises that typically fewer data can be generated via full-field simulations. Nevertheless, such sparse high quality data can be used to support the modeling with lower quality data. Concepts to incorporate such multi-fidelity data fusion approaches [67] in our SMTL model learning will be considered in the future.

Data availability

The data used to validate the SMTL approach is made available via the repository Fordatis at [https://fordatis.fraunhofer.de/handle/fordatis/204](https://fordatis.fraunhofer.de/handle/fordatis/204).
Author Contributions

TI developed and implemented the SMTL-framework as well as the optimization concept for materials design. LM set up the rolling texture generation model, implemented the properties calculation framework on the basis of the Taylor model and generated the used texture-property data. JD contributed theory and implementation of the crystallographic texture descriptor and distance function. NL supported in terms of machine learning. DH supported in terms of materials science with focus on the modelling of polycrystalline material behavior. All authors formulated the scientific challenge, contributed to the solution strategy and contributed to the discussion and the summary and outlook part of the paper.

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