



Dataset Documentation

Topology Optimization of the Base Cell of a Periodic Metamaterial

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1 Introduction

1.1 General Description

This work is part of the PhD thesis entitled *Análise de Sensibilidade de Variação Finita assistida por Redes Neurais Artificiais para Concepção de Metamateriais* (Finite Variation Sensitivity Analysis assisted by Artificial Neural Networks for Designing Metamaterials). The author Daniel Candeloro Cunha and his supervisor Professor Renato Pavanello are researchers at the Laboratory of Topology Optimization and Multiphysics Analysis, at the University of Campinas (Brazil).

The objective of the presented programs is to generate a dataset that will be used to train artificial neural networks. The purpose of such networks is to improve the performance of standard topology optimization programs, by reducing computational costs, making the procedures more stable, or more accurate.

The topology optimization of the base cell of a periodic metamaterial is considered. Two free parameters are used to define the inverse homogenization problem: the target Poisson's ratio and the minimal Young's modulus for the homogenized metamaterial. The dataset is generated by performing 18 382 optimizations, considering unique sets of these parameters.

All optimizations are performed through Sequential Integer Linear Programming (SILP). For each iteration of each case, all results are stored: topology vectors; sensitivity vectors; displacements vectors; homogenized Poisson's ratio values; homogenized Young's modulus values; volume fraction values. Also, metadata is stored with relevant information, for example, the corresponding input parameters of each result. This dataset occupies around 277 GB of disk.

- To collaborate or report bugs, you may contact the author: [cunhadc \[at\] unicamp \[dot\] br](mailto:cunhadc@unicamp.br)
- All codes and documentation are publicly available in the following github repository:
<https://github.com/Joquempo/Metamaterial-Dataset>

If you use the presented programs (or the data generated by it) in your work, the developer would be grateful if you would cite the indicated references. They are listed in the "CITEAS" file available in the github repository.

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1.2 Overview

In [section 2](#), the inverse homogenization problem is described. Then, the considered topology optimization method is presented.

In [section 3](#), a user guide is presented, explaining how to use the provided programs to generate the dataset. Then, each script is presented and briefly explained. Some validation procedures are described, the corresponding scripts are available in the github repository, but they are not shown in this document. Lastly, an unfixed bug, found late in development, is described (it is reported in the "Issues" tab of the github repository).

In [section 4](#), some samples are presented and discussed, to illustrate the data that composes the dataset.

In [section 5](#), a summary is presented with information about: the main parameters of the programs; the execution order of the scripts to generate the dataset; the data stored in the dataset.

2 Topology Optimization

2.1 Problem Description

The problem of designing the microstructure of a bidimensional isotropic mechanical metamaterial, in plane stress state, is considered. It corresponds to an inverse homogenization problem, in which we search for a topology of the microstructure that yields specified mechanical properties for the homogenized metamaterial. A hexagonal base cell with dihedral D_3 symmetry is used, which is a sufficient condition to obtain isotropic homogenized properties [1, 2, 3, 4]. **Figure 1** presents the base cell in the coordinate system (s_x, s_y) .

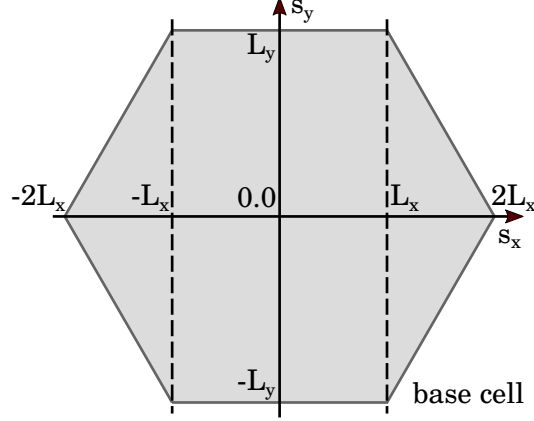


Figure 1: Coordinate system

The inverse homogenization problem can be solved through topology optimization. Since the optimization does not depend on the area of the base cell (V_Ω), or on the scale of the Young's modulus of the base material (\check{E}), unitary values are considered for both of them ($V_\Omega = 1.0 m^2$ and $\check{E} = 1.0 Pa$). Thus, $L_x = \left[\frac{1}{108}\right]^{\frac{1}{4}} m \approx 0.31 m$ and $L_y = \left[\frac{1}{12}\right]^{\frac{1}{4}} m \approx 0.54 m$.

Figure 2 presents the design domain. The six symmetries are indicated: the three rotations R_1 , R_2 and R_3 ; and the three reflections M_1 , M_2 and M_3 . The material distribution over the design domain determines the material distribution over the symmetric subdomains, through rotations and reflections.

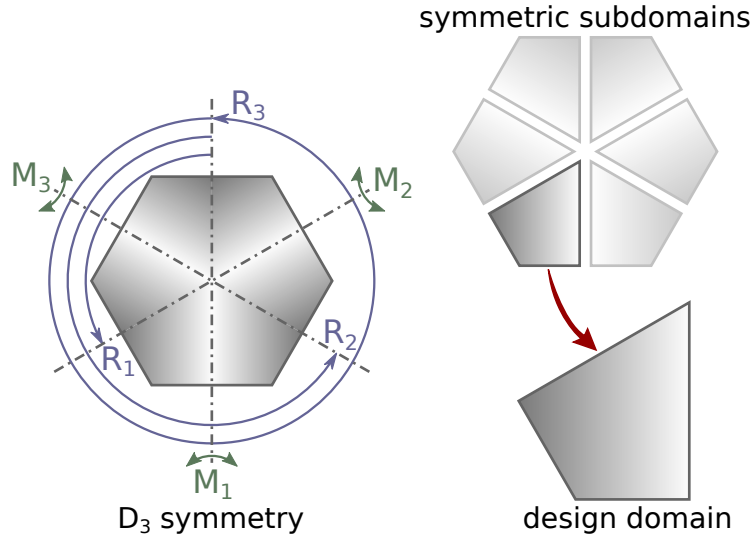


Figure 2: Design domain

For a fixed homogeneous and isotropic base material, of Poisson's ratio $\check{\nu} = 0.3$ and Young's modulus $\check{E} = 1.0 Pa$, the considered problem consists in finding a distribution of the base material over the design domain that yields a specified value (ν^*) for the homogenized Poisson's ratio ($\hat{\nu}$), while respecting a constraint of minimal homogenized Young's modulus ($\hat{E} \geq E_{\min}$).

In order to express this as an optimization problem, the structure is discretized in a mesh of finite elements,

then a computational homogenization [5, 6] is performed to obtain the required functions. The discretized structure is defined by the number of elements in each direction of the design domain, N_s . All elements have the same area, the length of their shorter sides is $e_x = \frac{L_x}{N_s}$ and the length of their longer sides is $e_y = \frac{L_y}{N_s}$. The number of design variables is given by $N_d = N_s^2$, and the number of quadrilateral finite elements in the cell is given by $N_t = 6 N_d$.

The design variables compose a density vector that fully describes the topology of the structure: $\mathbf{x} \in \{0, 1\}^{N_d}$. Each design variable defines the material distribution in six elements of the symmetric cell: when $x_i = 1$, all corresponding elements are solid and have the same stiffness of the base material; when $x_i = 0$, all corresponding elements are void and have nearly zero stiffness. The design variables, arranged in a matrix with proper neighborhood relations, are numbered from the leftmost column to the rightmost column and, in each column, they are numbered from bottom to top. The elements of the discretized cell are numbered from the center of the hexagon to the outer edges, following an anticlockwise spiral, as illustrated in Figure 3 for $N_s = 3$.

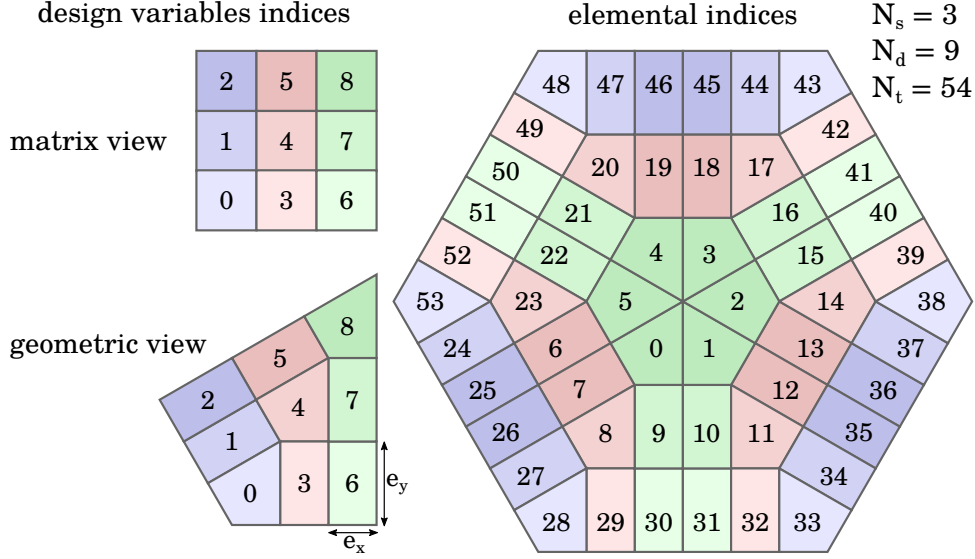


Figure 3: Elemental indices of the discretized cell

Figure 4 shows the nodal indices, which follow this same numbering rule.

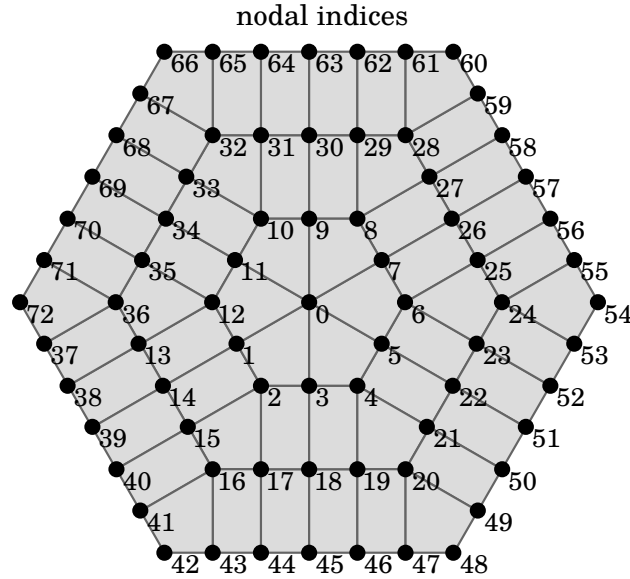


Figure 4: Nodal indices of the discretized cell

Bilinear quadrilateral elements in plane stress state are considered in the Finite Element Analysis (FEA). Small displacements and strains are considered, so linear assumptions are adopted. For each element, the local nodal indices are defined as shown in Figure 5, they are numbered anticlockwise. There are twelve different types of quadrilateral elements, according to their shape and rotation. Although some of them have the same stiffness

matrix, the twelve elemental stiffness matrices are independently computed, considering that the elements are solid, through Gaussian quadrature with 2×2 points.

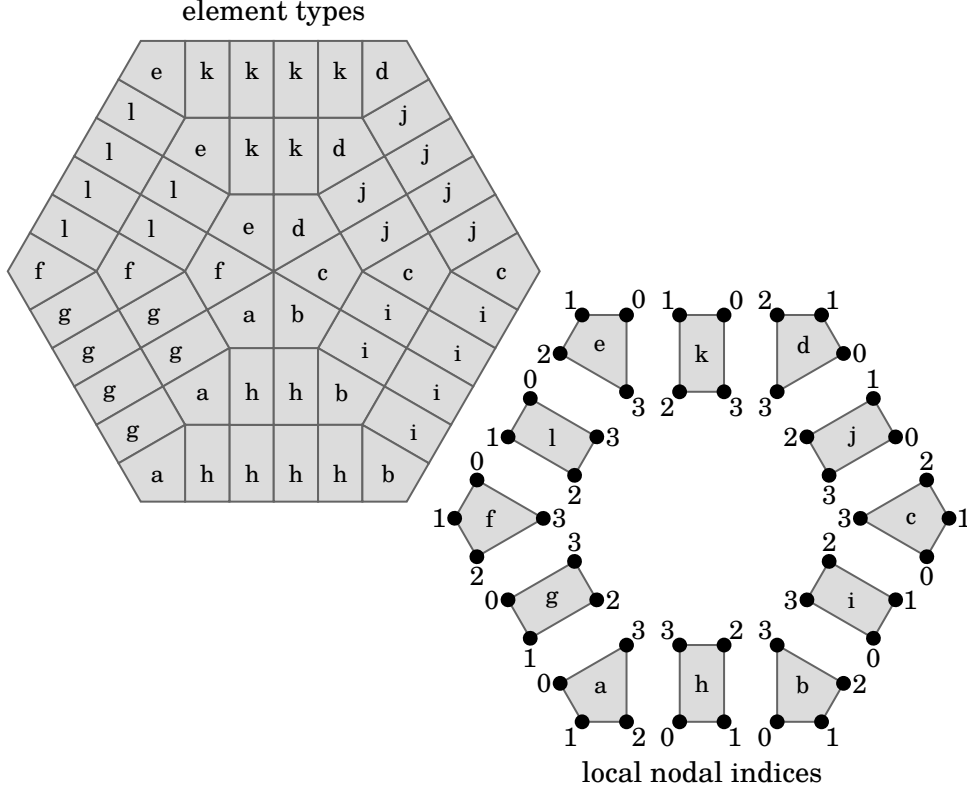


Figure 5: Local nodal indices of the bilinear quadrilateral elements

For simplicity, each sextuplet of symmetric elements (which share the same density value) can be understood as a single augmented element. Thus, for $x_i = 1$, the stiffness matrix of the i th augmented element is given by

$$\mathbf{K}_i^{[\text{aug}]} = \sum_{w=1}^6 \mathbf{K}_i^{[w]}, \quad (1)$$

where $\mathbf{K}_i^{[w]}$ is the stiffness matrix of the w th quadrilateral element corresponding to the i th design variable. The matrices of the quadrilateral elements are defined in the global system, so they assume zero values everywhere outside a small submatrix of dimensions 8×8 , and this assembly can be performed as a simple summation.

A soft-kill approach is adopted to prevent singularities throughout the optimization procedure, so a small stiffness is assigned to void elements according to a small soft-kill parameter (p_k). This means that the following base stiffness is assigned to the whole structure:

$$\mathbf{K}_0 = p_k \sum_{i=1}^{N_d} \mathbf{K}_i^{[\text{aug}]} . \quad (2)$$

The elemental variation matrix, corresponding to the variation applied to the global stiffness matrix when the state of the i th augmented element is switched, is defined as

$$\mathbf{K}_i = (1 - p_k) \mathbf{K}_i^{[\text{aug}]} . \quad (3)$$

Therefore, the global stiffness matrix of the base cell can be written as a function of \mathbf{x} :

$$\mathbf{K}(\mathbf{x}) = \mathbf{K}_0 + \sum_{i=1}^{N_d} x_i \mathbf{K}_i . \quad (4)$$

To perform the homogenization procedure [5, 6], three fixed macro-displacements are imposed. For a compact notation, they are grouped in a matrix:

$$\hat{\mathbf{U}} = \begin{bmatrix} \hat{\mathbf{u}}_{xx} & \hat{\mathbf{u}}_{yy} & \hat{\mathbf{u}}_{xy} \end{bmatrix}. \quad (5)$$

The total displacements is defined as

$$\mathbf{U}(\mathbf{x}) = \hat{\mathbf{U}} + \tilde{\mathbf{U}}(\mathbf{x}), \quad (6)$$

where $\tilde{\mathbf{U}}$ is a matrix storing the micro-displacement vectors. It can be written with respect to a reduced matrix $\check{\mathbf{U}}$ as

$$\tilde{\mathbf{U}}(\mathbf{x}) = \mathbf{P} \check{\mathbf{U}}(\mathbf{x}), \quad (7)$$

where \mathbf{P} is a known constant matrix used to impose the required periodicity constraints over the base cell of the metamaterial.

The reduced stiffness matrix and the reduced term for the right-hand side can then be defined as

$$\check{\mathbf{K}}(\mathbf{x}) = \mathbf{P}^T \mathbf{K}(\mathbf{x}) \mathbf{P} \text{ and } \check{\mathbf{F}}(\mathbf{x}) = -\mathbf{P}^T \mathbf{K}(\mathbf{x}) \hat{\mathbf{U}}, \quad (8)$$

resulting in a linear system with unique solution, from which $\check{\mathbf{U}}$ can be obtained:

$$\check{\mathbf{U}}(\mathbf{x}) = \left[\check{\mathbf{K}}(\mathbf{x}) \right]^{-1} \check{\mathbf{F}}(\mathbf{x}). \quad (9)$$

The elasticity matrix of the homogenized material is computed as follows:

$$\mathbf{C}(\mathbf{x}) = \begin{bmatrix} C_{00}(\mathbf{x}) & C_{01}(\mathbf{x}) & C_{02}(\mathbf{x}) \\ C_{10}(\mathbf{x}) & C_{11}(\mathbf{x}) & C_{12}(\mathbf{x}) \\ C_{20}(\mathbf{x}) & C_{21}(\mathbf{x}) & C_{22}(\mathbf{x}) \end{bmatrix} = \frac{1}{V_\Omega} [\mathbf{U}(\mathbf{x})]^T \mathbf{K}(\mathbf{x}) \mathbf{U}(\mathbf{x}) \quad (10)$$

Since isotropy is guaranteed, it must result in the following matrix:

$$\mathbf{C}(\mathbf{x}) = \frac{\hat{E}(\mathbf{x})}{1 - [\hat{\nu}(\mathbf{x})]^2} \begin{bmatrix} 1 & \hat{\nu}(\mathbf{x}) & 0 \\ \hat{\nu}(\mathbf{x}) & 1 & 0 \\ 0 & 0 & \frac{1-\hat{\nu}(\mathbf{x})}{2} \end{bmatrix}. \quad (11)$$

By comparing both expressions, the homogenized parameters can be obtained as

$$\hat{\nu}(\mathbf{x}) = 1 - \frac{2C_{22}(\mathbf{x})}{C_{00}(\mathbf{x})} \text{ and } \hat{E}(\mathbf{x}) = \frac{4C_{22}(\mathbf{x})[C_{00}(\mathbf{x}) - C_{22}(\mathbf{x})]}{C_{00}(\mathbf{x})}. \quad (12)$$

Besides the presented functions, volume and topology variation functions can be included in the problem in order to properly apply the considered optimization method. Since all elements have the same area, the number of solid augmented elements can be used as a volume function:

$$V(\mathbf{x}) = \|\mathbf{x}\|_1 = \sum_{i=1}^{N_d} x_i. \quad (13)$$

For a reference density vector $\bar{\mathbf{x}}$, the topology variation function can be defined as

$$D(\mathbf{x}) = \|\Delta \mathbf{x}\|_1 = \|\mathbf{x} - \bar{\mathbf{x}}\|_1 = \sum_{i=1}^{N_d} |x_i - \bar{x}_i|. \quad (14)$$

Both can be written as relative values, the volume fraction and topology variation fraction are given by:

$$V_f(\mathbf{x}) = \frac{V(\mathbf{x})}{N_d} \text{ and } D_f(\mathbf{x}) = \frac{D(\mathbf{x})}{N_d}. \quad (15)$$

Finally, for each pair of properties (ν^*, E_{\min}) , a topology optimization problem can be stated to solve the inverse homogenization problem.

$$\begin{aligned} \mathbf{x}^* &= \arg \min [\hat{\nu}(\mathbf{x}) - \nu^*]^2 \\ &\text{subject to} \\ &\hat{E}(\mathbf{x}) \geq E_{\min} \end{aligned} \quad (16)$$

Each optimization problem is then solved through SILP.

2.2 Sequential Integer Linear Programming (SILP)

In this approach, all functions have to be linearized around the current topology $\bar{\mathbf{x}}$ [7, 8, 9, 6]. In order to extract the linear component from a given function of binary variables, it is firstly written in the form

$$f(\mathbf{x}) = \alpha^{\langle 0 \rangle} + \sum_{j=1}^{N_d} \alpha^{\langle j \rangle} (\cdot)^j (\mathbf{x} - \bar{\mathbf{x}})^j, \quad (17)$$

where the j th-order tensor $(\mathbf{x} - \bar{\mathbf{x}})^j$ corresponds to the outer product between j vectors $(\mathbf{x} - \bar{\mathbf{x}})$ and the $(\cdot)^j$ -product represents the operation given by

$$\alpha^{\langle j \rangle} (\cdot)^j (\mathbf{x} - \bar{\mathbf{x}})^j = \sum_{i_1=1}^{N_d} \sum_{i_2=1}^{N_d} \dots \sum_{i_j=1}^{N_d} \alpha_{i_1 i_2 \dots i_j}^{\langle j \rangle} (x_{i_1} - \bar{x}_{i_1}) (x_{i_2} - \bar{x}_{i_2}) \dots (x_{i_j} - \bar{x}_{i_j}). \quad (18)$$

The scalar $\alpha^{\langle 0 \rangle}$ corresponds to $f(\bar{\mathbf{x}})$, $\alpha^{\langle 1 \rangle}$ is a vector of N_d entries corresponding to the variations of f when the state of a single element of $\bar{\mathbf{x}}$ is switched (from solid to void, or from void to solid), and, for $j > 1$, $\alpha^{\langle j \rangle}$ are strictly upper triangular tensors of order j , so $\alpha_{i_1 i_2 \dots i_j}$ only assumes non-zero values when $i_1 < i_2 < \dots < i_j$. The j -th order tensor, $\alpha^{\langle j \rangle}$, is related to the combined effect of simultaneously switching the state of j elements of $\bar{\mathbf{x}}$.

Thus, the linear truncation of $f(\mathbf{x})$, denoted by $f^{\text{lin}}(\mathbf{x})$, is given by

$$f^{\text{lin}}(\mathbf{x}) = \alpha^{\langle 0 \rangle} + \alpha^{\langle 1 \rangle} \cdot [\mathbf{x} - \bar{\mathbf{x}}] = f(\bar{\mathbf{x}}) + \alpha^{[f]} \cdot [\mathbf{x} - \bar{\mathbf{x}}]. \quad (19)$$

The vector $\alpha^{\langle 1 \rangle}$ is referred to as $\alpha^{[f]}$. It can be understood as the sensitivity of the linearized function with respect to each design variable, its entries are given by

$$\alpha_i^{[f]} = f(\bar{\mathbf{x}}, x_i = 1) - f(\bar{\mathbf{x}}, x_i = 0), \quad (20)$$

where the arguments $(\bar{\mathbf{x}}, x_i = 1)$ and $(\bar{\mathbf{x}}, x_i = 0)$ denote vectors that are equal to $\bar{\mathbf{x}}$ except at their i th term, which assumes the explicitly defined value.

The considered approach consists in solving a sequence of linearized subproblems, using the branch-and-bound method coupled with simplex algorithm to solve each linear integer subproblem [10, 11]. In the special case in which the Young's modulus is unconstrained ($E_{\min} = 0.0 Pa$), the BESO algorithm [12, 13] can be used instead, which is a much more efficient approach for problems with simple constraints.

The functions V_f and D_f are additively separable functions of binary variables, which means that they are already linear functions ($V_f = V_f^{\text{lin}}$ and $D_f = D_f^{\text{lin}}$) whose sensitivity values are given by

$$\alpha_i^{[V_f]} = \frac{1}{N_d} \text{ and } \alpha_i^{[D_f]} = \frac{1 - 2\bar{x}_i}{N_d}. \quad (21)$$

On the other hand, the considered objective function, $h_\nu(\mathbf{x}) = [\hat{\nu}(\mathbf{x}) - \nu^*]^2$, and constraint function, $g_E(\mathbf{x}) = E_{\min} - \hat{E}(\mathbf{x}) \leq 0$, are nonlinear. So their linearized versions should only be used as approximations for topologies reasonably close to $\bar{\mathbf{x}}$.

In order to improve the accuracy of the linearizations throughout the optimization procedure, a maximal topology variation (D_{\max}) is imposed in each iteration. An auxiliary parameter (η_E) is included to limit the

variation of \hat{E} . Moreover, a volume penalization parameter (β) is used to inhibit the appearance of disconnected solid islands in the structure.

Thus, to obtain the next topology of the iterative procedure ($\bar{\mathbf{x}}^{(k+1)}$), the functions are linearized around the current one ($\bar{\mathbf{x}}^{(k)}$) and the following subproblem is solved:

$$\begin{aligned} \bar{\mathbf{x}}^{(k+1)} = \arg \min & \quad h_{\nu}^{\text{lin}}(\mathbf{x}) + \beta V_f(\mathbf{x}) \\ \text{subject to} & \\ E_{\min} + \eta_E - \hat{E}^{\text{lin}}(\mathbf{x}) \leq 0 & \\ D_f(\mathbf{x}) \leq D_{\max} & \end{aligned} \quad (22)$$

The heuristic SILP approach consists in exploring the domain of feasible topologies by successively solving these linearized subproblems, starting from a given initial topology $\bar{\mathbf{x}}^{(0)}$. The stopping criterion is given in terms of a patience parameter (P): throughout the optimization procedure, the best topology thus far is stored, when P consecutive iterations are performed without obtaining a better topology, the procedure stops.

Usually, the most costly task of this algorithm is to perform the sensitivity analysis, that is, to compute $\alpha^{[h_{\nu}]}$ and $\alpha^{[E]}$. It is performed as follows.

The direct homogenization procedure is performed for the current topology ($\bar{\mathbf{x}}$), so all current matrices \mathbf{K} , $\tilde{\mathbf{U}}$, \mathbf{U} and \mathbf{C} are known, as well as the scalars $\tilde{\nu}$ and \tilde{E} . For a compact notation, the parameters $\bar{\gamma}$ and $\bar{\bar{\gamma}}$ are defined as

$$\bar{\gamma} = \frac{C_{22}}{C_{00}} \quad \text{and} \quad \bar{\bar{\gamma}} = \frac{C_{22} + \Delta C_{22}}{C_{00} + \Delta C_{00}}, \quad (23)$$

where ΔC_{00} and ΔC_{22} are diagonal terms of the matrix $\Delta \mathbf{C}$, which corresponds to the variation of \mathbf{C} when the state of the i th augmented is switched. Then, the variation of each elasticity property can be obtained as

$$\Delta \hat{\nu} = \frac{2[\bar{\gamma} \Delta C_{00} - \Delta C_{22}]}{C_{00} + \Delta C_{00}} \quad \text{and} \quad \Delta \hat{E} = 4[1 - \bar{\gamma}] \Delta C_{22} + 2 C_{22} \Delta \hat{\nu}. \quad (24)$$

Thus, the sensitivity values are obtained as

$$\alpha_i^{[h_{\nu}]} = \begin{cases} 2[\hat{\nu} - \nu^*] \Delta \hat{\nu} + \Delta \hat{\nu}^2, & \text{if } \bar{x}_i = 0, \\ -2[\hat{\nu} - \nu^*] \Delta \hat{\nu} + \Delta \hat{\nu}^2, & \text{if } \bar{x}_i = 1, \end{cases} \quad (25)$$

and

$$\alpha_i^{[E]} = \begin{cases} \Delta \hat{E}, & \text{if } \bar{x}_i = 0, \\ -\Delta \hat{E}, & \text{if } \bar{x}_i = 1, \end{cases} \quad (26)$$

The Woodbury Sensitivity (WS) approach [9, 6] can be used to compute the exact $\Delta \mathbf{C}$ for each augmented element. Firstly, the corresponding elemental variation matrix is factorized as

$$\mathbf{K}_i = \mathbf{H}_i \mathbf{H}_i^T, \quad (27)$$

where \mathbf{H}_i is a rectangular matrix with 30 columns at most, and with no more than 48 nonzero rows. Constrained matrices can be defined as

$$\tilde{\mathbf{H}}_i = \mathbf{P}^T \mathbf{H}_i \quad \text{and} \quad \tilde{\mathbf{K}}_i = \mathbf{P}^T \mathbf{K}_i \mathbf{P} = \tilde{\mathbf{H}}_i \tilde{\mathbf{H}}_i^T. \quad (28)$$

These are used to define the matrices

$$\mathbf{A}_i = \tilde{\mathbf{H}}_i^T \tilde{\mathbf{K}}^{-1} \tilde{\mathbf{H}}_i \quad \text{and} \quad \mathbf{V}_i = \mathbf{H}_i^T \mathbf{U} = \mathbf{H}_i^T \hat{\mathbf{U}} + \tilde{\mathbf{H}}_i^T \tilde{\mathbf{U}}. \quad (29)$$

Once \mathbf{A}_i and \mathbf{V}_i are known, the variation of \mathbf{C} can be obtained as

$$\Delta \mathbf{C} = \begin{cases} \frac{1}{V_\Omega} \mathbf{V}_i^T [\mathbf{I} + \mathbf{A}_i]^{-1} \mathbf{V}_i, & \text{if } \bar{x}_i = 0, \\ -\frac{1}{V_\Omega} \mathbf{V}_i^T [\mathbf{I} - \mathbf{A}_i]^{-1} \mathbf{V}_i, & \text{if } \bar{x}_i = 1. \end{cases} \quad (30)$$

Since the dimensions of \mathbf{A}_i are at most 30×30 , the inverse of $[\mathbf{I} \pm \mathbf{A}_i]$ can be easily computed. The Cholesky factorization is used to solve the linear systems from the homogenization procedure. Although this factorization can be reused to compute $\widetilde{\mathbf{K}}^{-1} \widetilde{\mathbf{H}}_i$, this has to be performed for each of the N_d augmented elements, thus, the computation of the exact sensitivity vectors is an expensive task.

Alternatively, approximated expressions can be used to reduce computational costs. The standard approximations are given by

$$\alpha_i^{[h\nu]} \approx 4 [\hat{\nu} - \nu^*] \left[\frac{\bar{\gamma} C_{00}^\delta - C_{22}^\delta}{C_{00}} \right] \quad (31)$$

and

$$\alpha_i^{[E]} \approx 4 [\bar{\gamma}^2 C_{00}^\delta + \hat{\nu} C_{22}^\delta], \quad (32)$$

where C_{00}^δ and C_{22}^δ are diagonal terms of the matrix

$$\mathbf{C}^\delta = \frac{1}{V_\Omega} \mathbf{U}^T \mathbf{K}_i \mathbf{U}. \quad (33)$$

However, by using the Conjugate Gradient Sensitivity (CGS) approach [9, 6], more accurate approximations can be obtained.

Jacobi preconditioning is used, so the preconditioner matrix \mathbf{M} corresponds to the diagonal of $[\widetilde{\mathbf{K}} + \Delta \widetilde{\mathbf{K}}]$, where $\Delta \widetilde{\mathbf{K}}$ is given by

$$\Delta \widetilde{\mathbf{K}} = \begin{cases} \widetilde{\mathbf{K}}_i, & \text{if } \bar{x}_i = 0, \\ -\widetilde{\mathbf{K}}_i, & \text{if } \bar{x}_i = 1. \end{cases} \quad (34)$$

The vector \mathbf{z}_h is used to compute the CGS approximations, when estimating ΔC_{00} , it is defined with respect to the first column of \mathbf{U} (\mathbf{u}_{xx}):

$$\mathbf{z}_h = \mathbf{P}^T \mathbf{K}_i \mathbf{u}_{xx}. \quad (35)$$

When estimating ΔC_{22} , \mathbf{z}_h is defined with respect to the third column of \mathbf{U} (\mathbf{u}_{xy}):

$$\mathbf{z}_h = \mathbf{P}^T \mathbf{K}_i \mathbf{u}_{xy}. \quad (36)$$

By computing, for each case, the vectors

$$\mathbf{z}_m = \mathbf{M}^{-1} \mathbf{z}_h \text{ and } \mathbf{z}_k = [\widetilde{\mathbf{K}} + \Delta \widetilde{\mathbf{K}}] \mathbf{z}_m, \quad (37)$$

and the coefficients

$$\omega_{hm} = \mathbf{z}_h^T \mathbf{z}_m, \quad \omega_{mk} = \mathbf{z}_m^T \mathbf{z}_k \text{ and } \phi_{m1} = \frac{\omega_{hm}}{\omega_{mk}}, \quad (38)$$

the CGS approximations with 1 step (CGS-1) can be obtained as

$$\Delta C_{00} \approx \begin{cases} C_{00}^\delta - \frac{1}{V_\Omega} [\phi_{m1} \omega_{hm}], & \text{if } \bar{x}_i = 0, \\ -C_{00}^\delta - \frac{1}{V_\Omega} [\phi_{m1} \omega_{hm}], & \text{if } \bar{x}_i = 1, \end{cases} \quad (39)$$

and

$$\Delta C_{22} \approx \begin{cases} C_{22}^\delta - \frac{1}{V_\Omega} [\phi_{m1} \omega_{hm}], & \text{if } \bar{x}_i = 0, \\ -C_{22}^\delta - \frac{1}{V_\Omega} [\phi_{m1} \omega_{hm}], & \text{if } \bar{x}_i = 1. \end{cases} \quad (40)$$

Using 2 steps, more accurate CGS expressions can be obtained. By computing, for each case, the vectors

$$\mathbf{z}_\eta = \mathbf{M}^{-1} \mathbf{z}_k \text{ and } \mathbf{z}_\xi = [\widetilde{\mathbf{K}} + \Delta \widetilde{\mathbf{K}}] \mathbf{z}_\eta, \quad (41)$$

and the coefficients

$$\omega_{k\eta} = \mathbf{z}_k^T \mathbf{z}_\eta, \quad \omega_{\eta\xi} = \mathbf{z}_\eta^T \mathbf{z}_\xi, \quad \phi_{m2} = \frac{\omega_{hm} \omega_{\eta\xi} - \omega_{mk} \omega_{k\eta}}{\omega_{mk} \omega_{\eta\xi} - \omega_{k\eta}^2} \text{ and } \phi_{\eta2} = \frac{\omega_{mk}^2 - \omega_{hm} \omega_{k\eta}}{\omega_{mk} \omega_{\eta\xi} - \omega_{k\eta}^2}, \quad (42)$$

the CGS-2 approximations can be obtained as

$$\Delta C_{00} \approx \begin{cases} C_{00}^\delta - \frac{1}{V_\Omega} [\phi_{m2} \omega_{hm} + \phi_{\eta2} \omega_{mk}], & \text{if } \bar{x}_i = 0, \\ -C_{00}^\delta - \frac{1}{V_\Omega} [\phi_{m2} \omega_{hm} + \phi_{\eta2} \omega_{mk}], & \text{if } \bar{x}_i = 1, \end{cases} \quad (43)$$

and

$$\Delta C_{22} \approx \begin{cases} C_{22}^\delta - \frac{1}{V_\Omega} [\phi_{m2} \omega_{hm} + \phi_{\eta2} \omega_{mk}], & \text{if } \bar{x}_i = 0, \\ -C_{22}^\delta - \frac{1}{V_\Omega} [\phi_{m2} \omega_{hm} + \phi_{\eta2} \omega_{mk}], & \text{if } \bar{x}_i = 1. \end{cases} \quad (44)$$

Instead of using the raw sensitivity vectors to linearize the functions and solve the optimization subproblem through a SILP algorithm, two procedures are included to improve the stability of the iterative procedure and the quality of the solutions. The sensitivity map is smoothed through a filtering procedure, this is done in order to deal with the checkerboard problem and mesh dependency [14, 15]. Moreover, a momentum method is included, so the previous values of the sensitivity vectors throughout the iterations (with proper weighting factors) are added to the value of the current sensitivity vectors [16]. This inhibits oscillations between consecutive iterations and favors a more extensive exploration of the domain of feasible topologies.

In order to properly filter the sensitivity map, the sensitivity values of the augmented elements are assigned to all corresponding symmetric quadrilateral elements in an extended mesh. The extended mesh includes parts of some neighboring periodic cells, and it is composed of $10 N_d$ elements, as illustrated in [Figure 6](#) for $N_s = 3$.

Here, a simple conical filter is considered, it is defined by its radius, r_s , which is a geometric parameter that indirectly controls the minimal thickness of the structural components, independently of the mesh.

Let $r(i, j)$ be defined as the distance between centers of the i -th and the j -th quadrilateral elements of the extended mesh. Then, the filter weights W_{ij} are given by

$$W_{ij} = \frac{\max(r_s - r(i, j), 0)}{\sum_{k=1}^{10 N_d} \max(r_s - r(i, k), 0)} \quad (45)$$

and the filtered sensitivity vector $[\boldsymbol{\alpha}^{[f]}]^{\text{fil}}$ is given by

$$[\alpha_i^{[f]}]^{\text{fil}} = \sum_{j=1}^{10 N_d} W_{ij} \alpha_j^{[f]}. \quad (46)$$

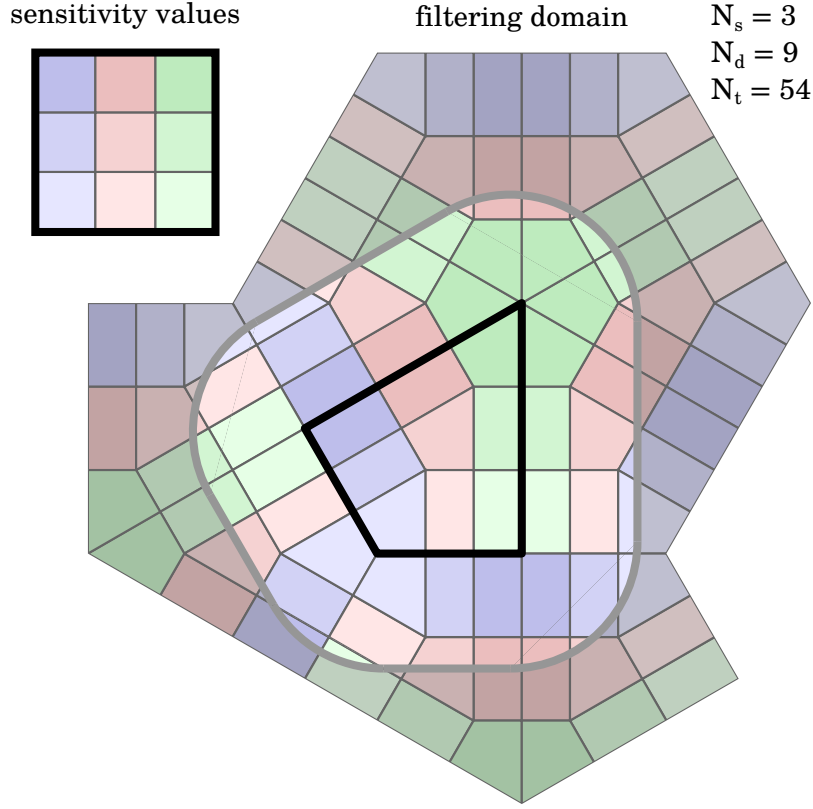


Figure 6: Extended mesh

Only elements within the range of the filter radius are considered when filtering the sensitivity value of each element. Furthest elements are disregarded since their contributions would be $\max(r_s - r(i, j), 0) = 0$. By definition, the conical filter weights decrease linearly with the distance from the central element. The filtering procedure is a linear transformation and the weights can be stored in a sparse matrix \mathbf{W} . Every row of \mathbf{W} adds up to 1, so there is no scaling factor.

For the momentum method, 25% is used for the objective function, and 0% (no momentum) is used for the constraint function. The sensitivity vector of the objective function is normalized so that eventual sensitivity peaks do not overly pollute the optimization process [17].

Thus, for the k -th iteration the normalized filtered vector $\tilde{\alpha}^{[h_\nu]}$ is given by

$$\tilde{\alpha}^{[h_\nu]} = \frac{[\alpha^{[h_\nu]}]^{\text{fil}}}{\|[\alpha^{[h_\nu]}]^{\text{fil}}\|_\infty} = \frac{[\alpha^{[h_\nu]}]^{\text{fil}}}{\max_i |[\alpha_i^{[h_\nu]}]^{\text{fil}}|}. \quad (47)$$

Then, the momentum is applied as follows:

$$[\alpha^{[h_\nu]}]^{\text{mom}(k)} = 0.75 \tilde{\alpha}^{[h_\nu](k)} + 0.25 [\tilde{\alpha}^{[h_\nu]}]^{\text{mom}(k-1)}. \quad (48)$$

Finally, the sensitivity vector used in the SILP algorithm is given by

$$[\tilde{\alpha}^{[h_\nu]}]^{\text{mom}} = \frac{[\alpha^{[h_\nu]}]^{\text{mom}}}{\|[\alpha^{[h_\nu]}]^{\text{mom}}\|_\infty} = \frac{[\alpha^{[h_\nu]}]^{\text{mom}}}{\max_i |[\alpha_i^{[h_\nu]}]^{\text{mom}}|}. \quad (49)$$

To perform this procedure consistently in the first iteration, it is defined that $[\tilde{\alpha}^{[h_\nu]}]^{\text{mom}(-1)} = \mathbf{0}$.

After solving the linearized subproblem, some additional procedures are performed. They correspond to reasonable small perturbations that are applied in each iteration to improve explorability and to bias the results towards simplified structures (with no disconnected solid islands, and with larger and smoother void cavities and solid components). In each iteration, an opening morphological operator is applied to the solution of the

linearized subproblem. Then, solid islands are identified. If the islands are composed by less than $N_d \times D_{\max}$ augmented elements, they are completely removed (turned into void); otherwise, the $N_d \times D_{\max}$ most insensitive augmented elements are removed, with respect to $[\alpha^{[E]}]^{\text{fil}}$. If the topology is kept unaltered after all these procedures (solving the linearized subproblem; applying the opening operator; and removing elements from solid islands), an erosion operator is applied to the topology. Lastly, if the constraint is not respected in the end of an iteration, the topology from the last iteration is recovered and a dilation operator is applied to it. When the optimization is concluded, remaining solid islands are removed from the final solution.

The morphological operators [18] are defined their radius r_m , which is a geometric parameter that controls the minimal thickness of the structural components. The operator weights B_{ij} are given by

$$B_{ij} = \begin{cases} 1, & \text{if } r(i, j) \leq r_m, \\ 0, & \text{if } r(i, j) > r_m. \end{cases} \quad (50)$$

As before, these distances are computed in the extended mesh, so that proper neighborhood relationships are considered for elements close to the boundaries of the design domain.

The eroded density vector is given by

$$[x_i]^{\text{ero}} = 1 - \max_j (B_{ij} [1 - x_j]), \quad (51)$$

the dilated density vector is given by

$$[x_i]^{\text{dil}} = \max_j (B_{ij} x_j), \quad (52)$$

and the opened density vector is given by

$$[x_i]^{\text{ope}} = \max_j (B_{ij} [x_j]^{\text{ero}}). \quad (53)$$

These operations can be efficiently performed by storing the required information in a sparse matrix \mathbf{B} .

In order to remove the disconnected structural components from the base cell, a depth-first search is performed to visit all solids connected to a reference element (arbitrarily chosen). Elements are considered to be connected if there is a path connecting them through a sequence of directly connected solid neighbors. Neighboring solid elements are considered to be directly connected if they have a common edge (elements that share a single node are not directly connected). To avoid identifying an island in place of the actual structure, depth-first searches are performed (altering the reference element) until the identified group of connected elements corresponds to at least 50% of the solids of the current topology. Then, all solid elements that are not in this group are eligible to be removed from the structure.

Once again, the extended mesh is used, so that proper neighborhood relationships are considered for paths going through the boundaries of the design domain.

3 Dataset Generation

3.1 Fixed Properties

The area of the base cell is set to $V_\Omega = 1.0 \text{ m}^2$, so the dimensions of the domain are $L_x = \left[\frac{1}{108}\right]^{\frac{1}{4}} \text{ m} \approx 0.31 \text{ m}$ and $L_y = \left[\frac{1}{12}\right]^{\frac{1}{4}} \text{ m} \approx 0.54 \text{ m}$. The number of elements in each direction of the design domain is set to $N_s = 32$, so the number of design variables is $N_d = 1024$, and the number of quadrilateral elements in the mesh is $N_t = 6144$. The sensitivity filter radius is set to $r_s = 0.024 \text{ m}$; and the radius used for the morphological operators is set to $r_m = 0.018 \text{ m}$.

The soft-kill parameter is set to $p_k = 1 \times 10^{-9}$; the volume penalization factor is set to $\beta = 0.05$; the maximal topology variation is set to $D_{\max} = 1.5625\% (1/64)$; η_E is defined in each iteration so that a maximal decrease of 0.05 Pa is imposed for the Young's modulus in each linearized subproblem; and the patience parameter used as stopping criterion is set to $P = 30$.

3.2 User Guide

All programs were developed in Python and Cython. Anaconda was used to manage packages through conda.

This short guide describes how to setup the conda environment, and how to properly execute the provided scripts in order to generate the datasets. Everything was developed in Linux (Ubuntu 20.04 LTS). There is a chance that other operating systems do not support some of the external packages used in the programs, if it is the case, users will have to adapt the codes according to their need.

3.2.1 Setup Conda Environment

To install Anaconda, go to <https://www.anaconda.com/distribution> and download the latest stable version. Alternatively, the Anaconda bash script may be downloaded using curl:

```
cd /tmp
curl -O https://repo.anaconda.com/archive/Anaconda3-2022.10-Linux-x86_64.sh
```

Then, run the bash script:

```
bash Anaconda3-2022.10-Linux-x86_64.sh
```

Follow the instructions to progress. Accept the license terms (if you agree with them) and specify your preferable location to install Anaconda. After the installation is finished, you will be asked if the installer should initialize Anaconda, write “yes”. Lastly, activates the installation:

```
source ~/.bashrc
```

Next, choose a location in your machine (<location_in_your_machine>) and download everything from the github repository (<https://github.com/Joquempo/Metamaterial-Dataset>). Finally, run the provided “metamaterial.sh” bash script in order to setup the conda environment:

```
cd <location_in_your_machine>/source
bash ./metamaterial.sh
```

This script will: update conda; create a new Python-3.8 environment named “metamaterial”; add the channel conda-forge; set channel_priority as strict; install numpy [19], scipy [20], matplotlib [21], cython [22], scikit-sparse [23] and pulp [24]; and build all Cython codes in “./cython” directory.

3.2.2 Generate Datasets

Before executing the scripts, be warned that the complete dataset occupies around **277 GB** of disk.

To generate the dataset, activate the metamaterial environment and go to the directory with the provided Python codes:

```
conda activate metamaterial
cd <location_in_your_machine>/source/python
```

Then, run the script “input_metamat.py”, which will create the “./input” directory and generate the input file “inputmat.npy”, it contains a set of 18382 unique pairs of parameters (ν^* , E_{\min}). This will occupy around 150 kB of disk.

```
python ./input_metamat.py
```

Although some procedures are embarrassingly parallelizable, it has been decided to keep the programs serialized and call multiple parallel executions, using multiple processors of the machine. Run the following command to list the processors in your machine. It is recommended to perform some tests to obtain the optimal number of parallel executions. In my machine, I used 4 processors, of indices 0, 1, 2 and 3 which are all different physical cores.

```
cat /proc/cpuinfo | egrep "processor|core id"
```

Then, go to “<location_in_your_machine>/source/python/SILP” and open the “basecell_silp.py” script in your preferable text editor. Redefine the values of the parameters “fid_ini” and “fid_lim” to select how many cases will be optimized. Create one copy of this script for each processor, with complementary values for the parameters “fid_ini” and “fid_lim”. For example: “fid_ini=0” and “fid_lim=700” in “basecell_silp_0.py”; “fid_ini=700” and “fid_lim=1400” in “basecell_silp_1.py”; “fid_ini=1400” and “fid_lim=2100” in “basecell_silp_2.py”; and “fid_ini=2100” and “fid_lim=2800” in “basecell_silp_3.py”. This will run the first 2800 cases using 4 processors.

The script prints information about which case is being optimized, so it is recommended that each script be executed in a different terminal window. Open the first terminal and execute the first script, setting it to the desired processor:

```
taskset -c 0 python ./SILP/basecell_silp_0.py
```

If it is the first time the program is being executed, the files “sfil_data.npy”, “sfil_row.npy”, “sfil_col.npy”, “mfil_data.npy”, “mfil_row.npy”, “mfil_col.npy” and “neighbors.npy” will be generated. This will occupy around 290 kB of disk. The file “neighbors.npy” contains a matrix of dimensions $N_d \times 4$ with the indices of all directly connected neighbors for each design variable. The other files contain the matrices (in COO format) of the sensitivity filter and morphological operators. It is recommended that no new parallel execution be started before all these files are generated. Then, open the next terminal and repeat for the next processor:

```
taskset -c 1 python ./SILP/basecell_silp_1.py
```

```
taskset -c 2 python ./SILP/basecell_silp_2.py
```

```
taskset -c 3 python ./SILP/basecell_silp_3.py
```

The “./SILP/output” directory will be created and each script will create a subfolder in it, to store the generated data. When the executions are concluded, verify the generated log files (in each subfolder), they present the input values corresponding to each problem, the obtained mechanical properties for the optimized metamaterials, and the execution time of the main tasks for each performed optimization.

Keep redefining the values of the parameters “fid_ini” and “fid_lim” (always with complementary values, so no redundant data is generated) and executing the scripts, until all 18 382 cases are optimized. Be warned that the WS expression (Equation 30), which is very costly, is computed in each iteration, so each optimization may take a few minutes (even for this coarse mesh). **Weeks may be necessary to generate the whole dataset, depending on your computer.**

After performing all 18 382 optimizations, run the script “generate_metamat.py” to conclude the generation of the dataset:

```
python ./generate_metamat.py
```

In the <location_in_your_machine>, this script will create the directory “./dataset/SILP”. Finally, it will relocate all generated data to this directory in an organized manner.

3.3 Implementation – Python

3.3.1 ./source/python/input_metamat.py

This script generates the input data for the inverse homogenization problems to be solved. For given properties of the base material (\tilde{E} and $\tilde{\nu}$), 18 382 unique pairs of prescribed properties (ν^* , E_{\min}) are defined and stored in a single file.

Firstly, the directory where the input data will be stored is created; and the parameters \tilde{E} and $\tilde{\nu}$ are defined, denoted by “Ey” and “nu”.

```

1 import os
2 import numpy as np
3 # check directories
4 if not os.path.exists('./input'):
5     os.mkdir('./input')
6 Ey = 1.00 # Young's modulus of the base material
7 nu = 0.30 # Poisson's ratio of the base material

```

Then, all considered values for the target Poisson’s ratio (ν^*) are generated and stored in the array “nuval_list”. Likewise, all considered values for the minimal Young’s modulus (E_{\min}) are generated and stored in the array “Eymin_list”.

The target Poisson’s ratio range from -1.00 to 1.00 in steps of 0.01 , but values close to the Poisson’s ratio of the base material (values in $]0.20, 0.40[$) are disregarded, which results in 182 unique values; the minimal Young’s modulus ranges from 0.0% to 50.0% in steps of 0.5% , relative to the Young’s modulus of the base material, which results in 101 unique values.

```

8  nuval_list = np.concatenate((np.linspace(-1.0,nu-0.1,1+int(100*(nu+0.9))),np.linspace(nu+0.1,1.0,1+int(100*(0.9-nu))))
9  nu_num = len(nuval_list)
10 Eymin_list = np.linspace(0,0.5*Ey,101)
11 Ey_num = len(Eymin_list)

```

Lastly, all values of target Poisson’s ratio are combined with all values of minimal Young’s modulus to generate all possible pairs (ν^*, E_{\min}), the total number of combinations is $18382 = 182 \times 101$. All pairs of properties are stored in the rows of the matrix “inputmat”. It is stored in disk (in 4-bytes format), in the input folder, named as “inputmat.npy”. It corresponds to around 145 kB of data.

```

12 print('generating input files')
13 total_num = nu_num*Ey_num
14 inputmat = np.ndarray([total_num,2],dtype=np.float32)
15 fid = 0
16 for knu in range(nu_num):
17     print(': {05d} / {05d} :'.format(knu*Ey_num+1,total_num))
18     for kEy in range(Ey_num):
19         input_num = knu*Ey_num + kEy
20         nuval = np.float32(nuval_list[knu]) # target Poisson's ratio
21         Eymin = np.float32(Eymin_list[kEy]) # minimal Young's modulus
22         inputmat[fid,0] = nuval
23         inputmat[fid,1] = Eymin
24         fid = fid + 1
25     print(': {05d} / {05d} :'.format(input_num+1,total_num))
26     np.save('./input/inputmat.npy',inputmat)
27     print('[ input file generated ]')
28     print('done!')

```

3.3.2 ./source/python/SILP/basecell_silp.py

This script performs the SILP optimization procedure for the selected input data. Two text files are written. The input-output log (“io_log.txt”) lists what this program generates as output data, presents each considered input data and the obtained mechanical properties of each optimized metamaterial, together with dates and times that inform when each optimization procedure was performed. The time log (“time_log.txt”) presents the execution times of the main tasks of the optimization procedures.

The output data is composed by numpy arrays with: the input files indices (“fid.npy”); the input data (“inp.npy”); the optimized topologies (“top_opt.npy”); the Poisson’s ratio of the optimized topology (“nu_opt.npy”); the Young’s modulus of the optimized topology (“Ey_opt.npy”); pointers relating each input with the corresponding iterations of the optimization processes (“ptr2opt.npy”); pointers relating each iteration of the optimization processes with the corresponding inputs (“ptr2inp.npy”); all generated topology vectors (“top.npy”); the displacements vectors corresponding to each generated topology (“dis_xx.npy”, “dis_yy.npy” and “dis_xy.npy”); the CGS-0 approximations for each topology (“dC00_0.npy”, “dC11_0.npy” and “dC22_0.npy”); the CGS-1 approximations (“dC00_1.npy”, “dC11_1.npy” and “dC22_1.npy”); the CGS-2 approximations (“dC00_2.npy”, “dC11_2.npy” and “dC22_2.npy”); the exact variations, obtained through the WS expression (“dC00_w.npy”, “dC11_w.npy” and “dC22_w.npy”); the Poisson’s ratio of each topology (“nu.npy”); the Young’s modulus of each topology (“Ey.npy”); the relative volume values of each topology (“vol.npy”); the execution times of the main tasks of the optimization processes, together with the number of performed iterations (“tim.npy”).

It should be noted that the arrays “dC00_0”, “dC00_1”, “dC00_2”, “dC00_w”, “dC11_0”, “dC11_1”, “dC11_2”, “dC11_w”, “dC22_0”, “dC22_1”, “dC22_2” and “dC22_w” store the **variations** of the diagonal terms of \mathbf{C} when the state of each augmented element is switched. They are sometimes referred to as sensitivity vectors because their absolute values correspond to the sensitivity values of these diagonal terms. However, to obtain proper sensitivity values, the signs of the values corresponding to solid augmented elements would have to be reversed.

Except for the density values, which are stored as single bits, all data is stored in 4-bytes format. For a more efficient storage of the generated data, each file stores the results from a group of 7 optimizations problems

(this value is defined through the parameter “noptf”). Thus, each “fid.npy” file contains 28 bytes of data; each “inp.npy” file contains 56 bytes of data; each “top_opt.npy” file contains 896 bytes of data; each “nu_opt.npy” file contains 28 bytes of data; each “Ey_opt.npy” file contains 28 bytes of data; each “ptr2opt.npy” file contains 32 bytes of data; and each “tim.npy” file contains 196 bytes of data. The other files depend on the number of iterations performed in each optimization process. Considering the average value of 74 iterations (so that each one of these lists has 75 entries), each “ptr2inp.npy” file would contain 2.1 kB of data; each “top.npy” file would contain 65.6 kB of data; each “dis_xx.npy” file would contain 25.4 MB of data; each “dis_yy.npy” file would contain 25.4 MB of data; each “dis_xy.npy” file would contain 25.4 MB of data; each “dC00_0.npy” would contain 2.1 MB of data; each “dC11_0.npy” would contain 2.1 MB of data; each “dC22_0.npy” would contain 2.1 MB of data; each “dC00_1.npy” would contain 2.1 MB of data; each “dC11_1.npy” would contain 2.1 MB of data; each “dC22_1.npy” would contain 2.1 MB of data; each “dC00_2.npy” would contain 2.1 MB of data; each “dC11_2.npy” would contain 2.1 MB of data; each “dC22_2.npy” would contain 2.1 MB of data; each “dC00_w.npy” would contain 2.1 MB of data; each “dC11_w.npy” would contain 2.1 MB of data; each “dC22_w.npy” would contain 2.1 MB of data; each “nu.npy” file would contain 2.1 kB of data; each “Ey.npy” file would contain 2.1 kB of data; and each “vol.npy” file would contain 2.1 kB of data. These add up to around 101 MB of data.

The 18382 optimization problems will generate 2626 of each of these files, which would result in around 260 GB of data. However, according to the Disk Usage Analyzer, a tool for analyzing disk usage for GNOME, the dataset occupies around **277 GB** of disk (7% more than the amount of useful data).

Firstly, the necessary modules are imported and all used-defined parameters are set.

```

1  import os, sys, gc
2  import numpy as np
3  from time import perf_counter
4  from datetime import datetime
5  from scipy.sparse import coo_matrix
6  from sksparse.cholmod import analyze
7  from mesh import get_mesh, get_fmsh
8  from elem import get_emat, get_augmat
9  from filters import get_sfil, get_mope
10 from rem_islands import visit, get_neighbors
11 from toptop import update, ws
12 from ilp_solver import solve_ILP, solve_BES0
13 sys.path.append('.../cython/')
14 from silp_sens import cgs
15 Ns = 32          # number of elements in each side of the design domain
16 Eyvar = 0.05     # maximal decrease in Young's modulus per iteration
17 nuvar = 2.0      # maximal variation in Poisson's ratio per iteration
18 Dmax = 0.015625  # maximal topology variation
19 rsen = 0.024     # sensitivity filter radius
20 rmor = 0.018     # morphology filter radius
21 patience = 30    # patience stop criterion
22 momentum = 0.25  # sensitivity momentum
23 beta = 0.05     # volume penalization factor
24 Ey = 1.00       # Young's modulus of the base material
25 nu = 0.30       # Poisson's ratio of the base material
26 pk = 1e-9       # soft-kill parameter
27 small = 1e-14   # small value to compare float numbers
28 noptf = 7       # number of optimizations to be stored in the same file
29 fid_ini = 0     # initial input index |run from input 0
30 fid_lim = 18382 # input index limit |up to input 18381

```

Some geometrical parameters are defined according to N_s (denoted by “Ns”), considering that the base cell is a regular hexagon of unitary area. The number of elements in the design domain (N_d) is stored in the variable “N”, and the number of quadrilateral elements in the mesh (N_t) is stored in the variable “Nt”. The variable “M” stores the number of nodes in the base cell, and the variable “G” stores the number of degrees of freedom of the unconstrained system. The mesh is generated by creating the matrix “coor” with the coordinates of each node of the mesh, and the matrix “inci” that relates each element to its nodes, this is performed by the function “mesh.get_mesh”. The array “etype” stores, for each quadrilateral element, the pointer to its corresponding stiffness matrix in “Ket” (defined in the next snippet of code), the matrix “sym” stores, for each design variable, the indices of its six corresponding quadrilateral elements. The python script “mesh.py” is detailed in a following section.

```

31 Lx = 1.0/(108**0.25) # design domain shorter side
32 Ly = np.sqrt(3)*Lx   # design domain longer side
33 Lex = Lx/Ns          # element shorter side
34 Ley = np.sqrt(3)*Lex # element longer side
35 N = Ns**2            # number of elements in the design domain
36 Nt = 6*N             # number of elements in the base cell
37 M = 1 + 6*Ns*(Ns+1)  # number of nodes in the base cell
38 G = 2*M              # number of degrees of freedom in the base cell

```

```

39 dXmax = int(round(N*Dmax)) # maximal topology variation (number of elements)
40 # Generate Mesh
41 coor, inci, etype, sym = get_mesh(Ns, Lex, Ley)

```

The function “elem.get_emat” is used to compute the all the elemental stiffness matrices, for the twelve considered types of bilinear quadrilateral elements, in plane stress state. They are stored in the tensor “Ket”, of dimensions $12 \times 8 \times 8$. Its vectorized data is stored in “Ketvec”. The stiffness variation matrices are stored in “dKe”, which correspond to the stiffness changes in the global matrix when the state of one element is switched (from solid to void, or from void to solid). The function “elem.get_augmat” is used to compute all the six different local stiffness variation matrices for the augmented elements, they are stored in the list “dKelist”. The array “aug_etype” stores, for each design variable, the pointer to its corresponding local matrix in “dKelist”. And the list “Hlist” stores the factorizations of each matrix in “dKelist” (Equation 27). The python script “elem.py” is detailed in a following section.

```

42 # Element Matrices (Quad4) - Plane Stress State
43 Ket = get_emat(Ey,nu)
44 Ketvec = np.ndarray((12,64))
45 dKe = np.ndarray((12,8,8))
46 for ek in range(12):
47     Ketvec[ek,:] = Ket[ek,:,:].ravel()
48     dKe[ek,:,:] = (1.0-pk)*Ket[ek,:,:] # stiffness variation of a topological change
49 # augmented element matrices (6xQuad4)
50 aug_etype, Hlist, dKelist = get_augmat(Ns,inci,etype,sym,dKe)

```

The initial topology is defined and stored in “x_init”. It is a mostly solid structure, with three rhombus-shaped cavities placed on different symmetry axes, each cavity is centered between the center of the hexagonal cell and one of its edges, each rhombus has a larger diagonal of $\frac{1}{16}$ of the height of the base cell and a smaller diagonal of $\frac{1}{16}$ of the side of the base cell. The symmetric density vector, which defines the density values for all the quadrilateral elements, is stored in “xt”.

```

51 # Initial Topology
52 x_init = np.ones(N,dtype=bool) # design variables
53 Ntotal = Ns//16
54 Nhole = Ntotal
55 while Nhole > 0:
56     x_init[(Ns-1-(Ntotal-Nhole))*Ns:(Ns-1-(Ntotal-Nhole))*Ns+(Ns//2+Nhole)] = False
57     Nhole = Nhole - 1
58 xt = np.ndarray((Nt),dtype=bool) # symmetric density vector
59 for k in range(N):
60     xt[sym[k,:]] = x_init[k]

```

If the file “sfil_data.npy” (“sfil_row.npy” and “sfil_col.npy” are not verified) is not in the “./source/python/SILP” directory, the sensitivity filter matrix is computed and stored in disk. If the file “mfil_data.npy” (“mfil_row.npy” and “mfil_col.npy” are not verified) does not exist in the directory, the morphological filter matrix is computed and stored in disk. Likewise, if the file “neighbors.npy” does not exist in the directory, the neighbors matrix is computed and stored in disk. If these files already exist, the corresponding matrices are loaded from disk. The function “mesh.get_fmsh” is used to generate the extended mesh, that is required to compute the sensitivity filter matrix, the morphological filter matrix and the neighbors matrix. The matrix “Q” is used to transform a vector of the design domain into a symmetric vector in the extended mesh. The function “filters.get_sfil” computes the sensitivity filter matrix, according to the defined radius r_s (denoted by “rsen”). The function “filters.get_mope” computes the morphological filter matrix, according to the defined radius r_m (denoted by “rmor”). The function “rem_islands.get_neighbors” computes the neighbors matrix, used to identify disconnected structural components. All these matrices occupy around 290 kB of disk. The python scripts “mesh.py”, “filters.py” and “rem_islands.py” are detailed in following sections.

```

61 # Generate Extended Mesh
62 if (not os.path.exists('./sfil_data.npy')) or (not os.path.exists('./mfil_data.npy')) or (not os.path.exists('./neighbors.npy')):
63     coor_lb, coor_bot, inci_lb, inci_bot, sym_lb, sym_bot = get_fmsh(Ns, Lx, Ly, Lex, Ley)
64 # Sensitivity and Morphology Filters Matrices
65 if os.path.exists('./sfil_data.npy') and os.path.exists('./mfil_data.npy'):
66     data = np.load('./sfil_data.npy')
67     row = np.load('./sfil_row.npy')
68     col = np.load('./sfil_col.npy')
69     Sf = coo_matrix((data,(row,col)),shape=(N,N))
70     Sf = Sf.tocsr()
71     data = np.load('./mfil_data.npy')
72     row = np.load('./mfil_row.npy')
73     col = np.load('./mfil_col.npy')

```

```

74     Mf = coo_matrix((data,(row,col)),shape=(N,N))
75     Mf = Mf.tocsr()
76 else:
77     # extended mesh
78     fcoor = np.vstack((coor,coor_lb,coor_bot))
79     finci = np.vstack((inci,inci_lb,inci_bot))
80     fsym = np.hstack((sym,sym_lb,sym_bot))
81     elepos = 0.25*fcoor[finci].sum(axis=1)
82     row = fsym.ravel('C')
83     col = np.repeat(np.arange(N),10)
84     data = np.ones(10*N)
85     Q = coo_matrix((data,(row,col)),shape=(10*N,N))
86     Q = Q.tocsc()
87     # sensitivity filter matrix
88     Sf = get_sfil(N,sym,elepos,Q,rsen)
89     Sfcoo = Sf.tocoo()
90     np.save('./sfil_data.npy',Sfcoo.data)
91     np.save('./sfil_row.npy',Sfcoo.row)
92     np.save('./sfil_col.npy',Sfcoo.col)
93     # morphology filter matrix
94     Mf = get_mope(N,sym,elepos,Q,rmor)
95     Mfcoo = Mf.tocoo()
96     np.save('./mfil_data.npy',Mfcoo.data)
97     np.save('./mfil_row.npy',Mfcoo.row)
98     np.save('./mfil_col.npy',Mfcoo.col)
99 # Get Neighbors
100 if os.path.exists('./neighbors.npy'):
101     neighbors = np.load('./neighbors.npy')
102 else:
103     neighbors = get_neighbors(Ns,inci,inci_lb,inci_bot,sym,sym_lb,sym_bot)
104     np.save('./neighbors.npy',neighbors)

```

The matrix of periodic boundary conditions is stored in “P”, it is used to impose zero-displacement at the node of index 0 (in the center of the hexagon), and to impose proper relations between displacements and loads of opposing edges. The macro-displacements vectors are computed and stored in the matrix “Uhat”.

```

105 # constraint matrix
106 Gb = 4*Ns
107 Gd = G - 2 - 6*Gb
108 Gr = Gd + 3*Gb - 2
109 ivec = np.arange(2,G)
110 j0 = np.arange(0,Gd)
111 j1 = np.arange(Gd,Gd+3*Gb-2)
112 v1=np.arange(Gd+Gb-2,Gd-1,-2)
113 v2=np.arange(Gd+Gb-1,Gd,-2)
114 j2=np.vstack((v1,v2)).ravel('F')
115 v1=np.arange(Gd+2*Gb-2,Gd+Gb-3,-2)
116 v2=np.arange(Gd+2*Gb-1,Gd+Gb-2,-2)
117 j3=np.vstack((v1,v2)).ravel('F')
118 v1=np.arange(Gd+3*Gb-4,Gd+2*Gb-3,-2)
119 v2=np.arange(Gd+3*Gb-3,Gd+2*Gb-2,-2)
120 j4=np.vstack((v1,v2)).ravel('F')
121 jvec = np.concatenate((j0,j1,j2,j3,j4))
122 avec = np.ones(G-2)
123 P = coo_matrix((avec,(ivec,jvec)),shape=(G,Gr)).tocsr()
124 # macro-strain tensors
125 eps_xx = np.array([[1,0],[0,0]])
126 eps_yy = np.array([[0,0],[0,1]])
127 eps_xy = np.array([[0,0.5],[0.5,0]])
128 # macro-displacements vectors
129 uhat_xx = np.ravel(coor @ eps_xx, 'C')
130 uhat_yy = np.ravel(coor @ eps_yy, 'C')
131 uhat_xy = np.ravel(coor @ eps_xy, 'C')
132 Uhat = np.vstack((uhat_xx,uhat_yy,uhat_xy)).T

```

If there is no input data, a standard input is created and stored in disk. The input matrix (denoted by “inputmat”) is loaded from disk. The directory where the output data will be stored is created. Both input-output log and time log files are opened to be written.

```

133 # check directories
134 if not os.path.exists('../input'):
135     os.mkdir('../input')
136 if not os.path.exists('../output'):
137     os.mkdir('../output')
138 if not os.path.exists('../output/run_{:05d}_{:05d}'.format(fid_ini,fid_lim-1)):
139     os.mkdir('../output/run_{:05d}_{:05d}'.format(fid_ini,fid_lim-1))
140 # check input
141 if not os.path.exists('../input/inputmat.npy'):
142     nuval = np.float32(0.00) # target Poisson's ratio
143     Eymin = np.float32(0.10) # minimal Young's modulus
144     inputmat = np.array([[nuval,Eymin]])
145     np.save('../input/inputmat.npy',inputmat)
146 # read input file

```

```

147 inputmat = np.load('../input/inputmat.npy')
148 # open log files
149 if not os.path.exists('../output/run_{:05d}_{:05d}/logs'.format(fid_ini,fid_lim-1)):
150     os.mkdir('../output/run_{:05d}_{:05d}/logs'.format(fid_ini,fid_lim-1))
151 iolog = open('../output/run_{:05d}_{:05d}/logs/io_log.txt'.format(fid_ini,fid_lim-1),'a')
152 tlog = open('../output/run_{:05d}_{:05d}/logs/time_log.txt'.format(fid_ini,fid_lim-1),'a')
153 iolog.truncate(0)
154 tlog.truncate(0)

```

The headers of the log files are written.

```

155 iolog.write('BASE CELL OPTIMIZATION (IO LOG)\n') # write in IO log
156 iolog.write('===== \n')
157 iolog.write(' OUTPUT :          input file id :          fid.npy          =\n')
158 iolog.write('----- :          input data :          inp.npy          =\n')
159 iolog.write('----- :          optimized topology :        top_opt.npy        =\n')
160 iolog.write('----- :          optimized Poisson\'s ratio :      nu_opt.npy        =\n')
161 iolog.write('----- :          optimized Young\'s modulus :      Ey_opt.npy        =\n')
162 iolog.write('----- :          pointer input > optimization :    ptr2opt.npy        =\n')
163 iolog.write('----- :          pointer optimization > input :    ptr2inp.npy        =\n')
164 iolog.write('----- :          topology vectors :          top.npy          =\n')
165 iolog.write('----- :          xx-displacements vectors :        dis_xx.npy        =\n')
166 iolog.write('----- :          yy-displacements vectors :        dis_yy.npy        =\n')
167 iolog.write('----- :          xy-displacements vectors :        dis_xy.npy        =\n')
168 iolog.write('----- :          dC00_CGS-0 sensitivity vectors :    dC00_0.npy        =\n')
169 iolog.write('----- :          dC00_CGS-1 sensitivity vectors :    dC00_1.npy        =\n')
170 iolog.write('----- :          dC00_CGS-2 sensitivity vectors :    dC00_2.npy        =\n')
171 iolog.write('----- :          dC00_WS sensitivity vectors :      dC00_w.npy        =\n')
172 iolog.write('----- :          dC11_CGS-0 sensitivity vectors :    dC11_0.npy        =\n')
173 iolog.write('----- :          dC11_CGS-1 sensitivity vectors :    dC11_1.npy        =\n')
174 iolog.write('----- :          dC11_CGS-2 sensitivity vectors :    dC11_2.npy        =\n')
175 iolog.write('----- :          dC11_WS sensitivity vectors :      dC11_w.npy        =\n')
176 iolog.write('----- :          dC22_CGS-0 sensitivity vectors :    dC22_0.npy        =\n')
177 iolog.write('----- :          dC22_CGS-1 sensitivity vectors :    dC22_1.npy        =\n')
178 iolog.write('----- :          dC22_CGS-2 sensitivity vectors :    dC22_2.npy        =\n')
179 iolog.write('----- :          dC22_WS sensitivity vectors :      dC22_w.npy        =\n')
180 iolog.write('----- :          Poisson\'s ratio array :          nu.npy          =\n')
181 iolog.write('----- :          Young\'s modulus array :          Ey.npy          =\n')
182 iolog.write('----- :          volume array :          vol.npy          =\n')
183 iolog.write('----- :          time array :          tim.npy          =\n')
184 iolog.write('===== \n')
185 iolog.write(' INPUT || NUVAL : EYMIN >> NUOPT : EYOPT ||          BEGIN :          END\n')
186 tlog.write('BASE CELL OPTIMIZATION (TIME LOG)\n') # write in time log
187 tlog.write('===== \n')
188 tlog.write(' INPUT || ( IT x ) :    M-ILP : M-SOLVER :    M-CGS :    M-WS : M-POST ||    TOTAL\n')

```

The analysis for the initial topology is performed. The matrix assembly is performed through the “scipy.sparse.coo_matrix” function. The variable “Kg_coo_init” is the COO (Coordinate list) unconstrained global stiffness matrix; “Kg_cgs” is the CSC (Compressed Sparse Column) unconstrained matrix; and “Kr” is the CSC constrained matrix. Since “scipy” automatically removes zero entries from the matrix after performing the constraining operations, a maneuver is performed to preserve the nonzero pattern of “Kr”. By preserving the nonzero pattern, the Cholesky factorization can be more efficiently performed throughout the optimization procedure. The variable “Fr” is the right-hand side of the linear system to be solved.

```

189 # Assembly
190 pen = np.ones(Nt)
191 pen[~xt] = pk
192 data = np.ndarray((64*Nt))
193 for et in range(Nt):
194     ek = etype[et]
195     data[64*et:64*et+64] = pen[et]*Ketvec[ek,:]
196 dof0 = 2*inci[:,0]
197 dof1 = dof0 + 1
198 dof2 = 2*inci[:,1]
199 dof3 = dof2 + 1
200 dof4 = 2*inci[:,2]
201 dof5 = dof4 + 1
202 dof6 = 2*inci[:,3]
203 dof7 = dof6 + 1
204 eledofs = np.array([dof0,dof1,dof2,dof3,dof4,dof5,dof6,dof7])
205 row = eledofs.repeat(8,axis=0).ravel('F')
206 col = eledofs.T.repeat(8,axis=0).ravel('C')
207 # stiffness matrix
208 Kg_coo_init = coo_matrix((data,(row,col)),shape=(G,G))
209 Kg_csc = Kg_coo_init.tocsc()
210 Kr = P.T @ Kg_csc @ P
211 # maneuver to fix the pattern of non-zero entries
212 Z_coo = coo_matrix((np.ones(64*Nt),(row,col)),shape=(G,G))
213 Z_csc = Z_coo.tocsc()
214 Zr = P.T @ Z_csc @ P
215 Zr.sort_indices()
216 shift = 10*np.amax(abs(Ket))

```

```

217 Kr = Kr + shift*Zr
218 Kr.sort_indices()
219 Kr.data = Kr.data - shift*Zr.data
220 # right-hand side
221 Fr = -P.T @ Kg_csc @ Uhat

```

The optimal fill-reducing permutation is computed for “Kr” using the “sksparse.cholmod.analyze” function. The reduced displacements matrix (\check{U}) is computed and stored in “Ur”. Then, the total displacements matrix (U) is stored in “Ug_init”.

```

222 # Solve System
223 factor = analyze(Kr)
224 factor.cholesky_inplace(Kr)
225 Ur = factor(Fr)
226 Ug_init = Uhat + P @ Ur

```

The elasticity matrix of the homogenized material, the corresponding mechanical properties and the volume fraction are computed for the initial topology.

```

227 # Effective Properties Matrix
228 Ch_init = Ug_init.T @ Kg_csc @ Ug_init
229 gamma_init = Ch_init[2,2]/Ch_init[0,0]
230 nuhat_init = 1-2*Ch_init[2,2]/Ch_init[0,0]
231 Eyhat_init = 4*Ch_init[2,2]*(Ch_init[0,0]-Ch_init[2,2])/Ch_init[0,0]
232 vol_init = sum(x_init)/N

```

The arrays that will store the sensitivity values of \hat{E} and h_ν are initialized. The sensitivity values of the diagonal terms of C are computed and stored. The function “silp_sens.cgs” is used to compute the CGS-0, CGS-1 and CGS-2 approximations, and the function “topopt.ws” is used to compute the exact sensitivity values through WS approach. The cython script “silp_sens.pyx” and the python script “topopt.py” are detailed in following sections.

```

233 s_Ey = np.ndarray((N))
234 s_obj = np.ndarray((N))
235 dC00_0_init = np.ndarray((N))
236 dC11_0_init = np.ndarray((N))
237 dC22_0_init = np.ndarray((N))
238 dC00_1_init = np.ndarray((N))
239 dC11_1_init = np.ndarray((N))
240 dC22_1_init = np.ndarray((N))
241 dC00_2_init = np.ndarray((N))
242 dC11_2_init = np.ndarray((N))
243 dC22_2_init = np.ndarray((N))
244 cgs(dC00_0_init,dC11_0_init,dC22_0_init,dC00_1_init,dC11_1_init,dC22_1_init,
245     dC00_2_init,dC11_2_init,dC22_2_init,x_init,N,sym,etype,aug_etype,inci,Ug_init,dKe,P,Kr,dKelist)
246 dC00_w_init, dC11_w_init, dC22_w_init = ws(x_init,aug_etype,sym,P,factor,inci,Ug_init,Hlist)

```

The loop to go through all the selected input values is started. The output data lists are initialized. A nested loop is started in order to store together the results of each block of “noptf” optimization problems.

```

247 file = 0 # file counter
248 fid = max([0,fid_ini])
249 while fid < min([fid_lim,inputmat.shape[0]]):
250     if not os.path.exists('./output/run_{:05d}_{:05d}/file_{:04d}'.format(fid_ini,fid_lim-1,file)):
251         os.mkdir('./output/run_{:05d}_{:05d}/file_{:04d}'.format(fid_ini,fid_lim-1,file))
252         list_fid = []
253         list_inp = []
254         list_top_opt = []
255         list_nu_opt = []
256         list_Ey_opt = []
257         list_ptr2opt = []
258         list_ptr2inp = []
259         list_top = []
260         list_dis_xx = []
261         list_dis_yy = []
262         list_dis_xy = []
263         list_dC00_0 = []
264         list_dC00_1 = []
265         list_dC00_2 = []
266         list_dC00_w = []
267         list_dC11_0 = []
268         list_dC11_1 = []
269         list_dC11_2 = []
270         list_dC11_w = []

```

```

271 list_dC22_0 = []
272 list_dC22_1 = []
273 list_dC22_2 = []
274 list_dC22_w = []
275 list_nu = []
276 list_Ey = []
277 list_vol = []
278 list_tim = []
279 ptr = 0 # pointer to input
280 for counter in range(noptf):
281     if fid >= min([fid_lim, inputmat.shape[0]]):
282         break
283     print('running : {05d} : setup'.format(fid))
284     inp_id = 'inp_{05d}'.format(fid)
285     iolog.write('> ' + inp_id + ' ||')
286     tlog.write('> ' + inp_id + ' ||')

```

The input values, in 4-bytes format, are read from “inputmat”. They are automatically cast to 8-bytes precision whenever they are used for comparisons or arithmetic operations. The input index is appended to “list_fid” and the input data is appended to “list_inp”. The input data is written in the input-output log. The topology vector, stiffness matrix, displacements matrix, elasticity matrix, mechanical properties, volume fraction, and sensitivity values of the diagonal terms of \mathbf{C} are recovered from the analysis performed for the initial topology.

```

287     nuval = inputmat[fid,0] # target Poisson's ratio
288     Eymin = inputmat[fid,1] # minimal Young's modulus
289     list_fid += [fid]
290     list_inp += [[nuval, Eymin]]
291     # write in log
292     iolog.write(' {5.2f} :'.format(nuval))
293     iolog.write(' {5.3f} >>'.format(Eymin))
294     begin = datetime.now().strftime(' %y/%m/%d-%H:%M:%S :')
295     # get initial data
296     x = x_init.copy()
297     Kg_coo = Kg_coo_init.copy()
298     Ug = Ug_init.copy()
299     Ch = Ch_init.copy()
300     gamma = gamma_init
301     nuhat = nuhat_init
302     Eyhat = Eyhat_init
303     vol = vol_init
304     dC00_0 = dC00_0_init.copy()
305     dC00_1 = dC00_1_init.copy()
306     dC00_2 = dC00_2_init.copy()
307     dC00_w = dC00_w_init.copy()
308     dC11_0 = dC11_0_init.copy()
309     dC11_1 = dC11_1_init.copy()
310     dC11_2 = dC11_2_init.copy()
311     dC11_w = dC11_w_init.copy()
312     dC22_0 = dC22_0_init.copy()
313     dC22_1 = dC22_1_init.copy()
314     dC22_2 = dC22_2_init.copy()
315     dC22_w = dC22_w_init.copy()

```

According to the maximal variation in Poisson’s ratio per iteration (“nuvar”), the parameter η_ν is computed. In the considered dataset, “nuvar” is set to 2.0, so η_ν is always 0.0 and no limitation imposed on the variation of $\hat{\nu}$. The function h_ν is computed and stored in “fnu_g”. The constraint function (that must be greater than or equal to zero) is computed and stored in “fEy”. According to the maximal decrease in Young’s modulus per iteration (“Eyvar”), to the constraint (“Eymin”) and to the Young’s modulus of the current topology (“Eyhat”), the parameter η_E is computed.

```

316     # target Poisson's ratio (limited variation)
317     if nuhat > nuval:
318         eta_nu = max([nuhat - nuval - nuvar, 0.0])
319     else:
320         eta_nu = min([nuhat - nuval + nuvar, 0.0])
321     nu_g = nuval + eta_nu
322     fnu_g = (nuhat - nu_g)**2
323     # minimal Young's modulus (limited variation)
324     fEy = Eyhat - Eymin
325     eta_Ey = max([fEy - Eyvar, 0.0])

```

The sensitivity analysis is performed for the initial topology. The raw sensitivity values of the \hat{E} and h_ν are stored in “s_Ey” and “s_obj”. The raw sensitivity values of the objective function (which includes the volume penalization) are stored in “raw_obj”. The objective function for the current topology is computed and stored in “obj”. The filtered sensitivity values (“fil_Ey” and “fil_obj”) are computed. The final sensitivity vector for

the objective function is stored in “mom_obj”, it is obtained by applying momentum from previous iterations (zero values in the first iteration) and by normalizing the vector with respect to its maximal absolute value.

```

326         # sensitivity analysis
327         for e in range(N):
328             ggamma = (Ch[2,2]+dC22_w[e])/(Ch[0,0]+dC00_w[e])
329             dnu = 2.0*(gamma*dC00_w[e]-dC22_w[e])/(Ch[0,0]+dC00_w[e])
330             dEy = 4.0*(1.0-ggamma)*dC22_w[e] + 2.0*Ch[2,2]*dnu
331             dobj = 2.0*(nuhat-nu_g)*dnu + dnu*dnu
332             if x[e]:
333                 s_Ey[e] = -dEy
334                 s_obj[e] = -dobj
335             else:
336                 s_Ey[e] = dEy
337                 s_obj[e] = dobj
338             raw_obj = s_obj + beta/N
339             obj = fnu_g + beta*vol
340             fil_Ey = Sf @ s_Ey
341             fil_obj = Sf @ raw_obj
342             mom_obj = np.zeros(N)
343             mom_obj = momentum*mom_obj + (1.0-momentum)*fil_obj/max(abs(fil_obj))
344             mom_obj = mom_obj/max(abs(mom_obj))

```

The list of pointers “list_ptr2opt” appends the value that relates the current input to the index of the current optimization in the block of “noptf” processes. The list of pointers “list_ptr2inp” appends the value that relates the index of the current iteration in the block of “noptf” optimization processes to the current input. The current topology, displacements vectors, sensitivity values of the diagonal terms of \mathbf{C} , mechanical properties and volume fraction are appended to their corresponding lists. For the current optimization, the best topology obtained thus far is stored in “top_opt”, the values of its mechanical properties and objective function are stored in “nu_opt”, “Ey_opt” and “obj_opt”.

```

345         # store data
346         size_list = len(list_ptr2inp)
347         list_ptr2opt += [size_list]
348         list_ptr2inp += [ptr]
349         list_top     += [x.copy()]
350         list_dis_xx  += [Ug[:,0].copy()]
351         list_dis_yy  += [Ug[:,1].copy()]
352         list_dis_xy  += [Ug[:,2].copy()]
353         list_dC00_0  += [dC00_0.copy()]
354         list_dC00_1  += [dC00_1.copy()]
355         list_dC00_2  += [dC00_2.copy()]
356         list_dC00_w  += [dC00_w.copy()]
357         list_dC11_0  += [dC11_0.copy()]
358         list_dC11_1  += [dC11_1.copy()]
359         list_dC11_2  += [dC11_2.copy()]
360         list_dC11_w  += [dC11_w.copy()]
361         list_dC22_0  += [dC22_0.copy()]
362         list_dC22_1  += [dC22_1.copy()]
363         list_dC22_2  += [dC22_2.copy()]
364         list_dC22_w  += [dC22_w.copy()]
365         list_nu      += [nuhat]
366         list_Ey      += [Eyhat]
367         list_vol     += [vol]
368         # optimized topology thus far
369         top_opt = x.copy()
370         nu_opt = nuhat
371         Ey_opt = Eyhat
372         obj_opt = obj

```

The time array is initialized and the optimization loop is started. Elements that are certain to keep their state are disregarded in the current iteration, the array “selection” stores to the boolean mask that identifies the elements that may change their state. If there is no Young’s modulus constraint, the “ilp_solver.solve_BESO” function is used to solve the linear subproblem with the BESO algorithm. Otherwise, the “ilp_solver.solve_ILP” function is used to solve it with branch-and-bound simplex. After solving the linear subproblem, the opening morphological operator is applied to the topology vector. Then, disconnected solid elements are removed. If the topology is kept unaltered, the erosion operator is applied to it. The time to perform these procedures is stored in “time_array[0]”. The python script “ilp_solver.py” is detailed in a following section.

```

373         time_array = np.zeros(7) # initialize time array
374         keep_going = True
375         waiting = 0
376         it = 0
377         while keep_going:
378             it = it + 1
379             print('running : :05d : :5d'.format(fid,it))

```

```

380 # solve ILP
381 t0 = perf_counter()
382 selection = (x & (mom_obj>-small)) | ((~x) & (mom_obj<small)) | (x & (fil_Ey<small)) | ((~x) & (fil_Ey>-small))
383 Nsel = sum(selection)
384 y = x.copy()
385 if Nsel > 0:
386     if (Eymin + eta_Ey) < small:
387         ysel = solve_BESO(Nsel,x[selection],mom_obj[selection],dXmax)
388         y[selection] = ysel
389     else:
390         ysel = solve_ILP(Nsel,x[selection],mom_obj[selection],fil_Ey[selection],fEy,eta_Ey,dXmax,sense_h='G')
391         y[selection] = ysel
392 # open operator (erode + dilate)
393 y[Mf[-y,:].indices] = False
394 y[Mf[y,:].indices] = True
395 # remove islands
396 voly = sum(y)/N
397 continent = np.zeros(N,dtype=bool)
398 continent_vol = 0.0
399 for e in (list(range(0,Ns))+list(range(Ns,N,Ns))):
400     if y[e] and (not continent[e]):
401         continent = np.zeros(N,dtype=bool)
402         visit(e,y,continent,neighbors)
403         continent_vol = sum(continent)/N
404     if continent_vol > 0.50*voly:
405         break
406 if continent_vol > 0.50*voly:
407     islands = np.argwhere(y!=continent).ravel()
408     if len(islands) > dXmax:
409         sortedargs = np.argsort(abs(fil_Ey[islands]))
410         y[islands[sortedargs[:dXmax]]] = False
411     else:
412         y[islands] = False
413 # erode if nothing has been changed
414 if all(x==y):
415     print('--- erode ---')
416     y[Mf[-y,:].indices] = False
417 t1 = perf_counter()
418 time_array[0] += (t1-t0)

```

The function “topopt.update” is used to update the topology vector, the stiffness matrix and the displacements matrix. The new mechanical properties, volume fraction, constraint function and objective function are computed. If the new topology breaks the Young’s modulus constraint, the previous topology is recovered, the dilation operator is applied to it, and everything is computed again for this new topology. The time to perform these procedures is stored in “time_array[1]”. The python script “topopt.py” is detailed in a following section.

```

418 # update topology
419 t0 = perf_counter()
420 if any(x!=y):
421     elist = list(np.argwhere(x!=y)[: ,0])
422     Ug, Kr = update(x,etype,sym,pk,Ketvec,P,Kg_coo,Zr,shift,Uhat,factor,elist)
423 # compute homogenized properties
424 Kg_csc = Kg_coo.tocsc()
425 Ch = Ug.T @ Kg_csc @ Ug
426 gamma = Ch[2,2]/Ch[0,0]
427 nuhat = 1-2*Ch[2,2]/Ch[0,0]
428 Eyhat = 4*Ch[2,2]*(Ch[0,0]-Ch[2,2])/Ch[0,0]
429 if nuhat > nuval:
430     eta_nu = max([nuhat - nuval - nuvar, 0.0])
431 else:
432     eta_nu = min([nuhat - nuval + nuvar, 0.0])
433 nu_g = nuval + eta_nu
434 fnu_g = (nuhat-nu_g)**2
435 vol = sum(x)/N
436 obj = fnu_g + beta*vol
437 fEy_test = Eyhat-Eymin
438 eta_Ey_test = max([fEy_test - Eyvar, 0.0])
439 # go back to last topology and dilate if constraint is broken
440 if fEy_test < 0.0:
441     # go back to last topology
442     update(x,etype,sym,pk,Ketvec,P,Kg_coo,Zr,shift,Uhat,factor,elist,solve_sys=False)
443     # dilate
444     print('--- dilate ---')
445     y = x.copy()
446     y[Mf[y,:].indices] = True
447     elist = list(np.argwhere(x!=y)[: ,0])
448     # compute homogenized properties
449     Ug, Kr = update(x,etype,sym,pk,Ketvec,P,Kg_coo,Zr,shift,Uhat,factor,elist)
450     Kg_csc = Kg_coo.tocsc()
451     Ch = Ug.T @ Kg_csc @ Ug
452     gamma = Ch[2,2]/Ch[0,0]
453     nuhat = 1-2*Ch[2,2]/Ch[0,0]
454     Eyhat = 4*Ch[2,2]*(Ch[0,0]-Ch[2,2])/Ch[0,0]
455     if nuhat > nuval:

```



```

456         eta_nu = max([nuhat - nuval - nuvar, 0.0])
457     else:
458         eta_nu = min([nuhat - nuval + nuvar, 0.0])
459     nu_g = nuval + eta_nu
460     fnu_g = (nuhat-nu_g)**2
461     vol = sum(x)/N
462     obj = fnu_g + beta*vol
463     fEy = Eyhat-Eymin
464     eta_Ey = max([fEy - Eyvar, 0.0])
465 else:
466     fEy = fEy_test
467     eta_Ey = eta_Ey_test
468     t1 = perf_counter()
469     time_array[1] += (t1-t0)

```

The CGS and WS sensitivity values for the diagonal terms of \mathbf{C} are computed with the functions “silp_sens.cgs” and “topopt.ws”. The time to perform the CGS analysis is stored in “time_array[2]” and the time to perform the WS analysis is stored in “time_array[3]”. The cython script “silp_sens.pyx” and the python script “topopt.py” are detailed in following sections.

```

470         # CGS analysis for dCh
471         t0 = perf_counter()
472         cgs(dC00_0,dC11_0,dC22_0,dC00_1,dC11_1,dC22_1,dC00_2,dC11_2,dC22_2,
473             x,N,sym,etype,aug_etype,inci,Ug,dKe,P,Kr,dKelist)
474         t1 = perf_counter()
475         time_array[2] += (t1-t0)
476         # WS analysis for dCh
477         t0 = perf_counter()
478         dC00_w, dC11_w, dC22_w = ws(x,aug_etype,sym,P,factor,inci,Ug,Hlist)
479         t1 = perf_counter()
480         time_array[3] += (t1-t0)

```

The sensitivity analysis is performed for the current topology. The data of the current iteration is appended to the output lists. If the Young’s modulus constraint is respected, and if the Poisson’s ratio is better than the best result obtained thus far, the best topology obtained thus far is updated. In order to improve explorability, if the objective function has decreased, even if the Poisson’s ratio is worse than the best result obtained thus far, the counter used to check convergence (“waiting”) is set to zero. If the limit number of iterations without improvements (“patience”) has been achieved, the boolean variable “keep_going” is set to *False*, which will end the optimization loop. The time to perform these procedures is stored in “time_array[4]”.

```

481         # post-procedures
482         t0 = perf_counter()
483         # sensitivity analysis for obj and Ey
484         for e in range(N):
485             gamma = (Ch[2,2]+dC22_w[e])/(Ch[0,0]+dC00_w[e])
486             dnu = 2.0*(gamma*dC00_w[e]-dC22_w[e])/(Ch[0,0]+dC00_w[e])
487             dEy = 4.0*(1.0-ggamma)*dC22_w[e] + 2.0*Ch[2,2]*dnu
488             dobj = 2.0*(nuhat-nu_g)*dnu + dnu*dnu
489             if x[e]:
490                 s_Ey[e] = -dEy
491                 s_obj[e] = -dobj
492             else:
493                 s_Ey[e] = dEy
494                 s_obj[e] = dobj
495         raw_obj = s_obj + beta/N
496         obj = fnu_g + beta*vol
497         fil_Ey = Sf @ s_Ey
498         fil_obj = Sf @ raw_obj
499         mom_obj = momentum*mom_obj + (1.0-momentum)*fil_obj/max(abs(fil_obj))
500         mom_obj = mom_obj/max(abs(mom_obj))
501         # store data
502         list_ptr2inp += [ptr]
503         list_top += [x.copy()]
504         list_dis_xx += [Ug[:,0].copy()]
505         list_dis_yy += [Ug[:,1].copy()]
506         list_dis_xy += [Ug[:,2].copy()]
507         list_dC00_0 += [dC00_0.copy()]
508         list_dC00_1 += [dC00_1.copy()]
509         list_dC00_2 += [dC00_2.copy()]
510         list_dC00_w += [dC00_w.copy()]
511         list_dC11_0 += [dC11_0.copy()]
512         list_dC11_1 += [dC11_1.copy()]
513         list_dC11_2 += [dC11_2.copy()]
514         list_dC11_w += [dC11_w.copy()]
515         list_dC22_0 += [dC22_0.copy()]
516         list_dC22_1 += [dC22_1.copy()]
517         list_dC22_2 += [dC22_2.copy()]
518         list_dC22_w += [dC22_w.copy()]
519         list_nu += [nuhat]
520         list_Ey += [Eyhat]

```

```

521     list_vol     += [vol]
522     # stopping criterion
523     if ((obj<(1.0-small)*obj_opt) or (abs(nuhat-nuval)<(1.0-small)*abs(nu_opt-nuval))) and (Eyhat>Eymin-small):
524         waiting = 0
525         obj_opt = obj
526         if (abs(nuhat-nuval)<(1.0-small)*abs(nu_opt-nuval)):
527             # update optimized topology
528             top_opt = x.copy()
529             nu_opt  = nuhat
530             Ey_opt  = Eyhat
531     else:
532         waiting += 1
533         # check convergence
534         if waiting == patience:
535             keep_going = False
536         t1 = perf_counter()
537         time_array[4] += (t1-t0)

```

After concluding the current optimization process, all remaining disconnected solid elements are removed from the optimized topology. This produces the unfixed bug presented in [subsection 3.7](#).

```

538     # remove remaining islands from optimized topology
539     y = top_opt.copy()
540     voly = sum(y)/N
541     continent = np.zeros(N,dtype=bool)
542     continent_vol = 0.0
543     for e in (list(range(0,Ns))+list(range(Ns,N,Ns))):
544         if y[e] and (not continent[e]):
545             continent = np.zeros(N,dtype=bool)
546             visit(e,y,continent,neighbors)
547             continent_vol = sum(continent)/N
548             if continent_vol > 0.50*voly:
549                 break
550     if continent_vol > 0.50*voly:
551         islands = np.argwhere(y!=continent).ravel()
552         y[islands] = False
553     if any(top_opt!=y):
554         if any(x!=y):
555             elist = list(np.argwhere(x!=y)[: ,0])
556             Ug, Kr = update(x,etype,sym,pk,Ketvec,P,Kg_coo,Zr,shift,Uhat,factor,elist)
557             Kg_csc = Kg_coo.tocsc()
558             Ch = Ug.T @ Kg_csc @ Ug
559             gamma = Ch[2,2]/Ch[0,0]
560             nuhat = 1-2*Ch[2,2]/Ch[0,0]
561             Eyhat = 4*Ch[2,2]*(Ch[0,0]-Ch[2,2])/Ch[0,0]
562             top_opt = x.copy()
563             nu_opt = nuhat
564             Ey_opt = Eyhat

```

The total time to perform the current optimization process is stored in “time_array[5]”. The time array is updated to store the average times of the tasks performed in the optimization loop. The number of iterations performed in the current optimization process is stored in “time_array[6]”. The execution times are written in the time log. The mechanical properties of the optimized topology are written in the input-output log. The optimized topology and the corresponding mechanical properties are appended to “list_top_opt”, “list_nu_opt” and “list_Ey_opt”. The “time_array” is appended to “list_tim”. The pointer variable “ptr” and the input index variable “fid” are updated so the optimization process for the next input can start.

```

564     # write in log
565     tlog.write(' ({:4d} x )'.format(it))
566     time_array[5] = sum(time_array[:5])
567     time_array[5] = time_array[5]/it
568     time_array[6] = (1+small)*it
569     tlog.write(' {:6.3f} s : {:6.3f} s : {:6.3f} s : {:6.3f} s : {:6.3f} s ||'.format(
570         time_array[0],time_array[1],time_array[2],time_array[3],time_array[4]))
571     tlog.write(' {:7.1f} s\n'.format(time_array[5]))
572     iolog.write(' {:5.2f} '.format(nu_opt))
573     iolog.write(' {:5.3f} ||'.format(Ey_opt))
574     iolog.write(begin)
575     iolog.write(datetime.now().strftime(' %y/%m/%d-%H:%M:%S\n'))
576     # store data
577     list_top_opt += [top_opt.copy()]
578     list_nu_opt  += [nu_opt]
579     list_Ey_opt  += [Ey_opt]
580     list_tim     += [time_array.copy()]
581     # update pointer
582     ptr += 1
583     # prepare to open next input file
584     fid += 1

```

After performing “noptf” optimization processes, the corresponding output data is written in disk. The last value of “list_ptr2opt” is appended. Then, each output list is saved in an independent file. Afterward, the

output variables are deleted and “gc.collect()” is called to ensure that the RAM be freed. The file counter “file” is updated so the next block of “noptf” optimization processes can start.

```

585         size_list = len(list_ptr2inp)
586         list_ptr2opt += [size_list]
587         # save files
588         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/fid.npy'.format(
589             fid_ini,fid_lim-1,file),np.array(list_fid,dtype=np.uint32))
590         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/inp.npy'.format(
591             fid_ini,fid_lim-1,file),np.array(list_inp,dtype=np.float32))
592         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/top_opt.npy'.format(
593             fid_ini,fid_lim-1,file),np.packbits(np.array(list_top_opt),axis=1))
594         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/nu_opt.npy'.format(
595             fid_ini,fid_lim-1,file),np.array(list_nu_opt,dtype=np.float32))
596         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/Ey_opt.npy'.format(
597             fid_ini,fid_lim-1,file),np.array(list_Ey_opt,dtype=np.float32))
598         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/ptr2opt.npy'.format(
599             fid_ini,fid_lim-1,file),np.array(list_ptr2opt,dtype=np.uint32))
600         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/ptr2inp.npy'.format(
601             fid_ini,fid_lim-1,file),np.array(list_ptr2inp,dtype=np.uint32))
602         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/top.npy'.format(
603             fid_ini,fid_lim-1,file),np.packbits(np.array(list_top),axis=1))
604         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dis_xx.npy'.format(
605             fid_ini,fid_lim-1,file),np.array(list_dis_xx,dtype=np.float32))
606         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dis_yy.npy'.format(
607             fid_ini,fid_lim-1,file),np.array(list_dis_yy,dtype=np.float32))
608         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dis_xy.npy'.format(
609             fid_ini,fid_lim-1,file),np.array(list_dis_xy,dtype=np.float32))
610         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC00_0.npy'.format(
611             fid_ini,fid_lim-1,file),np.array(list_dC00_0,dtype=np.float32))
612         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC00_1.npy'.format(
613             fid_ini,fid_lim-1,file),np.array(list_dC00_1,dtype=np.float32))
614         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC00_2.npy'.format(
615             fid_ini,fid_lim-1,file),np.array(list_dC00_2,dtype=np.float32))
616         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC00_w.npy'.format(
617             fid_ini,fid_lim-1,file),np.array(list_dC00_w,dtype=np.float32))
618         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC11_0.npy'.format(
619             fid_ini,fid_lim-1,file),np.array(list_dC11_0,dtype=np.float32))
620         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC11_1.npy'.format(
621             fid_ini,fid_lim-1,file),np.array(list_dC11_1,dtype=np.float32))
622         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC11_2.npy'.format(
623             fid_ini,fid_lim-1,file),np.array(list_dC11_2,dtype=np.float32))
624         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC11_w.npy'.format(
625             fid_ini,fid_lim-1,file),np.array(list_dC11_w,dtype=np.float32))
626         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC22_0.npy'.format(
627             fid_ini,fid_lim-1,file),np.array(list_dC22_0,dtype=np.float32))
628         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC22_1.npy'.format(
629             fid_ini,fid_lim-1,file),np.array(list_dC22_1,dtype=np.float32))
630         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC22_2.npy'.format(
631             fid_ini,fid_lim-1,file),np.array(list_dC22_2,dtype=np.float32))
632         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/dC22_w.npy'.format(
633             fid_ini,fid_lim-1,file),np.array(list_dC22_w,dtype=np.float32))
634         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/nu.npy'.format(
635             fid_ini,fid_lim-1,file),np.array(list_nu,dtype=np.float32))
636         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/Ey.npy'.format(
637             fid_ini,fid_lim-1,file),np.array(list_Ey,dtype=np.float32))
638         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/vol.npy'.format(
639             fid_ini,fid_lim-1,file),np.array(list_vol,dtype=np.float32))
640         np.save('./output/run_{:05d}_{:05d}/file_{:04d}/tim.npy'.format(
641             fid_ini,fid_lim-1,file),np.array(list_tim,dtype=np.float32))
642         del list_fid, list_inp, list_top_opt, list_nu_opt, list_Ey_opt, list_ptr2opt, list_ptr2inp, list_top
643         del list_dis_xx, list_dis_yy, list_dis_xy, list_dC00_0, list_dC00_1, list_dC00_2, list_dC00_w
644         del list_dC11_0, list_dC11_1, list_dC11_2, list_dC11_w, list_dC22_0, list_dC22_1, list_dC22_2, list_dC22_w
645         del list_nu, list_Ey, list_vol, list_tim
646         gc.collect()
647         # prepare to write next output file
648         file += 1

```

When all selected optimization processes are done, the log files are closed and the program terminates.

```

649         iolog.close()
650         tlog.close()
651         print('done!')

```

3.3.3 ./source/python/mesh.py

This script generates the hexagonal mesh for the base cell, and the extended mesh used to perform filtering operations and to identify directly connected neighbors. Firstly, the numpy module is imported.

```

1     import numpy as np

```

The function “get_mesh” is defined. It generates the matrix “coor” with the coordinates of each node; the matrix “inci” with the nodes corresponding to each quadrilateral element; the array “etype” with the types of each quadrilateral element; and the matrix “sym” which relates each design variable with its six corresponding quadrilateral elements.

```

2 def get_mesh(Ns, Lex, Ley):
3     N = Ns**2
4     Nt = 6*N
5     M = 1 + 6*Ns*(Ns+1)
6     # coordinates matrix
7     coor = np.ndarray((M,2))
8     coor[0,:] = [0.0,0.0]
9     for k in range(Ns):
10        num = 1 + 6*k*(k+1)
11        pts = np.array([[-2*k*Lex-1.5*Lex, -0.5*Ley],
12                        [-(k+1)*Lex, -(k+1)*Ley],
13                        [-k*Lex, -(k+1)*Ley],
14                        [(k+1)*Lex, -(k+1)*Ley],
15                        [k*Lex + 1.5*Lex, -k*Ley-0.5*Ley],
16                        [2*(k+1)*Lex, 0.0],
17                        [2*k*Lex+1.5*Lex, 0.5*Ley],
18                        [(k+1)*Lex, (k+1)*Ley],
19                        [k*Lex, (k+1)*Ley],
20                        [-(k+1)*Lex, (k+1)*Ley],
21                        [-k*Lex-1.5*Lex, k*Ley+0.5*Ley],
22                        [-2*(k+1)*Lex, 0.0]])
23        for kk in range(6):
24            coor[num+2*kk*(k+1):num+2*(kk+1)*(k+1),0] = np.linspace(pts[2*kk,0],pts[2*kk+1,0],2*(k+1))
25            coor[num+2*kk*(k+1):num+2*(kk+1)*(k+1),1] = np.linspace(pts[2*kk,1],pts[2*kk+1,1],2*(k+1))
26        # incidence matrix
27        inci = np.ndarray((Nt,4),dtype=np.uint32)
28        etype = np.ndarray((Nt),dtype=np.uint8)
29        for k in range(Ns):
30            for kk in range(6):
31                ek = 6*(k**2) + 2*(kk+1)*k + kk
32                etype[ek] = kk
33                inci[ek,0] = (2*kk+1) + (14+2*kk)*k + 6*k*(k-1)
34                inci[ek,1] = (2*kk+2) + (14+2*kk)*k + 6*k*(k-1)
35                if kk == 5:
36                    inci[ek,2] = (2*kk+3) + (14+2*kk)*(k-1) + 6*(k-1)*(k-2)
37                else:
38                    inci[ek,2] = (2*kk+3) + (14+2*kk)*k + 6*k*(k-1)
39                    inci[ek,3] = (2*kk+2) + (14+2*kk)*(k-1) + 6*(k-1)*(k-2)
40        for k in range(1,Ns):
41            for kk in range(6):
42                ek = 6 + 18*(k-1) + 6*(k-2)*(k-1) + (2*k+1)*kk
43                etype[ek:ek+2*k] = 6 + kk
44                kkk = (kk-1)%6
45                if kkk == 5:
46                    inci[ek:ek+2*k,0] = (2*kkk+3) + (14+2*kkk)*(k-1) + 6*(k-1)*(k-2)
47                    inci[ek:ek+2*k,1] = (2*kkk+4) + (14+2*kkk)*(k-1) + 6*(k-1)*(k-2)
48                    inci[ek:ek+2*k,2] = (2*kkk+3) + (14+2*kkk)*(k-2) + 6*(k-2)*(k-3)
49                    inci[ek:ek+2*k,3] = (2*kkk+2) + (14+2*kkk)*(k-2) + 6*(k-2)*(k-3)
50                else:
51                    inci[ek:ek+2*k,0] = (2*kkk+3) + (14+2*kkk)*k + 6*k*(k-1)
52                    inci[ek:ek+2*k,1] = (2*kkk+4) + (14+2*kkk)*k + 6*k*(k-1)
53                    inci[ek:ek+2*k,2] = (2*kkk+3) + (14+2*kkk)*(k-1) + 6*(k-1)*(k-2)
54                    inci[ek:ek+2*k,3] = (2*kkk+2) + (14+2*kkk)*(k-1) + 6*(k-1)*(k-2)
55                    inci[ek:ek+2*k,:] = inci[ek:ek+2*k,:] + np.arange(2*k).reshape(2*k,1)
56                ek = 6 + 18*(k-1) + 6*(k-2)*(k-1)
57                inci[ek,3] = 12 + 24*(k-1) + 6*(k-1)*(k-2)
58        # D3-symmetry map
59        sym = np.ndarray((N,6),dtype=np.uint32)
60        for k in range(Ns):
61            for kk in range(6):
62                ek = 6*(Ns-k-1)**2 + 2*(kk+1)*(Ns-k-1) + kk
63                kkk = 1-2*(kk%2)
64                sym[k*(Ns+1)::Ns,kk] = np.array(range(ek,ek+kk*(Ns-k),kkk))
65                if kk == 5:
66                    ek = 6*(Ns-k-2)**2 + 2*(kk+1)*(Ns-k-2) + kk
67                    sym[k*(Ns+1)+1:(k+1)*Ns,kk] = np.array(range(ek-kkk,ek-kkk*(Ns-k),-kkk))
68        return coor, inci, etype, sym

```

The function “get_fmesh” is defined. It generates the matrix “coor_lb” with the coordinates of the nodes of the cell located at the left-bottom of the main cell; the matrix “coor_bot” with the coordinates of the nodes of the cell located at the bottom of the main cell; the matrix “inci_lb” with the nodes corresponding to each quadrilateral element of the left-bottom cell; the matrix “inci_bot” with the nodes corresponding to each quadrilateral element of the bottom cell; the matrix “sym_lb” which relates each design variable with its two corresponding quadrilateral elements of the left-bottom cell; and the matrix “sym_bot” which relates each design variable with its two corresponding quadrilateral elements of the bottom cell.

```

69 def get_fmesh(Ns, Lx, Ly, Lex, Ley):
70     N = Ns**2
71     M = 1 + 6*Ns*(Ns+1)
72     # left-bottom cell
73     Mlb = 2*N + Ns
74     coor_lb = np.ndarray((Mlb,2))
75     coor_lb[0,:] = [0.0,0.0]
76     for k in range(Ns-1):
77         num = 1 + 5*k + 2*(k-1)*k
78         pts = np.array([[ 1.5*(k+1)*Lex , -0.5*(k+1)*Ley],
79                        [ 2*(k+1)*Lex , 0.0],
80                        [ 2*k*Lex+1.5*Lex , 0.5*Ley],
81                        [ (k+1)*Lex , (k+1)*Ley],
82                        [ k*Lex , (k+1)*Ley],
83                        [ 0.0 , (k+1)*Ley]])
84         coor_lb[num : num+ k+2,0] = np.linspace(pts[0,0],pts[1,0], k+2)
85         coor_lb[num : num+ k+2,1] = np.linspace(pts[0,1],pts[1,1], k+2)
86         coor_lb[num+ k+2:num+3*k+4,0] = np.linspace(pts[2,0],pts[3,0],2*k+2)
87         coor_lb[num+ k+2:num+3*k+4,1] = np.linspace(pts[2,1],pts[3,1],2*k+2)
88         coor_lb[num+3*k+4:num+4*k+5,0] = np.linspace(pts[4,0],pts[5,0], k+1)
89         coor_lb[num+3*k+4:num+4*k+5,1] = np.linspace(pts[4,1],pts[5,1], k+1)
90     k = Ns-1
91     num = 1 + 5*k + 2*(k-1)*k
92     pts = np.array([[ 1.5*(k+1)*Lex , -0.5*(k+1)*Ley],
93                    [ 2*k*Lex+1.5*Lex , -0.5*Ley],
94                    [ k*Lex , (k+1)*Ley],
95                    [ 0.0 , (k+1)*Ley]])
96     coor_lb[num : num+ k+1,0] = np.linspace(pts[0,0],pts[1,0], k+1)
97     coor_lb[num : num+ k+1,1] = np.linspace(pts[0,1],pts[1,1], k+1)
98     coor_lb[num+k+1:num+2*k+2,0] = np.linspace(pts[2,0],pts[3,0], k+1)
99     coor_lb[num+k+1:num+2*k+2,1] = np.linspace(pts[2,1],pts[3,1], k+1)
100    coor_lb = coor_lb - np.array([3*Lx,Ly])
101    inci_lb = np.ndarray((2*N,4),dtype=np.uint32)
102    etype_lb = np.ndarray((2*N),dtype=np.int8)
103    for k in range(Ns-1):
104        for kk in range(2):
105            ek = 2*((k+kk)**2) + k - kk*(2*k+1)
106            etype_lb[ek] = 2+kk
107            id1 = 1 + 5*(k+kk) + 2*(k-1+kk)*(k+kk) + (k+1) - 2*kk*(k+1) - kk
108            id3 = 1 + 5*(k-1+kk) + 2*(k-2+kk)*(k-1+kk) + k - 2*kk*k - kk
109            inci_lb[ek,0] = id1-1
110            inci_lb[ek,1] = id1
111            inci_lb[ek,2] = id1+1
112            inci_lb[ek,3] = id3
113    k = Ns-1
114    for kk in range(2):
115        ek = 2*((k+kk)**2) + k - kk*(2*k+1)
116        etype_lb[ek] = 2+kk
117        if kk == 0:
118            inci_lb[ek,0] = 2*(k**2) + 4*k + 1
119            inci_lb[ek,1] = 6*k*(k+1) + 2*Ns - M
120            inci_lb[ek,2] = 6*k*(k+1) + 2*Ns - 1 - M
121            inci_lb[ek,3] = 2*(k**2)
122        else:
123            inci_lb[ek,0] = 6*k*(k+1) + 1 - M
124            inci_lb[ek,1] = 6*(k+1)*(k+2) - M
125            inci_lb[ek,2] = 2*(k**2) + 4*k + 3
126            inci_lb[ek,3] = 2*(k**2) + 2*k
127    for k in range(1,Ns-1):
128        kkk = np.tile(np.arange(4*k), (2,1))
129        kkk[0, k:] = kkk[0, k:] + 2
130        kkk[0,3*k:] = kkk[0,3*k:] + 2
131        for kk in range(3):
132            ek = 2*(k**2) + kk + k*((kk*(kk+1))/2)
133            etype_lb[ek:ek+((2-kk)*kk+1)*k] = 8 + kk
134            n0 = 1 + 5*k + 2*(k-1)*k
135            n1 = n0 + 1
136            n2 = 2 + 5*(k-1) + 2*(k-2)*(k-1)
137            n3 = n2 - 1
138            inci_lb[ek:ek+((2-kk)*kk+1)*k,0] = n0 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
139            inci_lb[ek:ek+((2-kk)*kk+1)*k,1] = n1 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
140            inci_lb[ek:ek+((2-kk)*kk+1)*k,2] = n2 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
141            inci_lb[ek:ek+((2-kk)*kk+1)*k,3] = n3 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
142    k = Ns-1
143    kkk = np.tile(np.arange(4*k), (2,1))
144    kkk[0,k:3*k] = np.arange(0,-2*Ns+2,-1)
145    kkk[0,3*k:] = kkk[0,3*k:] - 2*k + 1
146    for kk in range(3):
147        ek = 2*(k**2) + kk + k*((kk*(kk+1))/2)
148        etype_lb[ek:ek+((2-kk)*kk+1)*k] = 8 + kk
149        if kk == 1:
150            n0 = 1 + 6*k*(k+1) + 2*(Ns-1) - M
151            n1 = n0 - 1
152        else:
153            n0 = 1 + 5*k + 2*(k-1)*k
154            n1 = n0 + 1
155            n2 = 2 + 5*(k-1) + 2*(k-2)*(k-1)
156            n3 = n2 - 1
157            inci_lb[ek:ek+((2-kk)*kk+1)*k,0] = n0 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
158            inci_lb[ek:ek+((2-kk)*kk+1)*k,1] = n1 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
159            inci_lb[ek:ek+((2-kk)*kk+1)*k,2] = n2 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]

```

```

160     inci_lb[ek:ek+((2-kk)*kk+1)*k,3] = n3 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
161 inci_lb = inci_lb + M
162 sym_lb = np.ndarray((N,2),dtype=np.uint32)
163 for k in range(Ns):
164     for kk in range(2):
165         ek = 2*((k+kk)**2) + k - kk*(2*k+1)
166         sym_lb[N-1-k*(Ns+1):N-k*Ns,kk] = np.arange(ek,ek+(2*kk-1)*(k+1),2*kk-1)
167 for k in range(Ns-1):
168     for kk in range(2):
169         sym_lb[(k+1)*Ns+k:Ns,kk] = np.arange(sym_lb[k*Ns+k,kk]+1-2*kk,sym_lb[k*Ns+k,kk]+(1-2*kk)*(Ns-k),1-2*kk)
170 sym_lb = sym_lb + 6*N
171 # bottom cell
172 Mbot = 2*N
173 coor_bot = np.ndarray((Mbot,2))
174 coor_bot[0,:] = [0.0,0.0]
175 for k in range(Ns-1):
176     num = 1 + 5*k + 2*(k-1)*k
177     pts = np.array([[ 1.5*(k+1)*Lex , 0.5*(k+1)*Ley],
178                    [ (k+1)*Lex , (k+1)*Ley],
179                    [ k*Lex , (k+1)*Ley],
180                    [ -(k+1)*Lex , (k+1)*Ley],
181                    [ -k*Lex-1.5*Lex , k*Ley+0.5*Ley],
182                    [ -1.5*(k+1)*Lex , 0.5*(k+1)*Ley]])
183     coor_bot[num : num+ k+2,0] = np.linspace(pts[0,0],pts[1,0], k+2)
184     coor_bot[num : num+ k+2,1] = np.linspace(pts[0,1],pts[1,1], k+2)
185     coor_bot[num+ k+2:num+3*k+4,0] = np.linspace(pts[2,0],pts[3,0],2*k+2)
186     coor_bot[num+ k+2:num+3*k+4,1] = np.linspace(pts[2,1],pts[3,1],2*k+2)
187     coor_bot[num+3*k+4:num+4*k+5,0] = np.linspace(pts[4,0],pts[5,0], k+1)
188     coor_bot[num+3*k+4:num+4*k+5,1] = np.linspace(pts[4,1],pts[5,1], k+1)
189 k = Ns-1
190 num = 1 + 5*k + 2*(k-1)*k
191 pts = np.array([[ 1.5*(k+1)*Lex , 0.5*(k+1)*Ley],
192                [ k*Lex + 1.5*Lex , k*Ley+0.5*Ley]])
193 coor_bot[num:num+k+1,0] = np.linspace(pts[0,0],pts[1,0], k+1)
194 coor_bot[num:num+k+1,1] = np.linspace(pts[0,1],pts[1,1], k+1)
195 coor_bot = coor_bot - np.array([0.0,2*Ly])
196 inci_bot = np.ndarray((2*N,4),dtype=np.uint32)
197 etype_bot = np.ndarray((2*N),dtype=np.int8)
198 for k in range(Ns-1):
199     for kk in range(2):
200         ek = 2*((k+kk)**2) + k - kk*(2*k+1)
201         etype_bot[ek] = 3+kk
202         id1 = 1 + 5*(k+kk) + 2*(k-1+kk)*(k+kk) + (k+1) - 2*kk*(k+1) - kk
203         id3 = 1 + 5*(k-1+kk) + 2*(k-2+kk)*(k-1+kk) + k - 2*kk*k - kk
204         inci_bot[ek,0] = id1-1
205         inci_bot[ek,1] = id1
206         inci_bot[ek,2] = id1+1
207         inci_bot[ek,3] = id3
208 k = Ns-1
209 for kk in range(2):
210     ek = 2*((k+kk)**2) + k - kk*(2*k+1)
211     etype_bot[ek] = 3+kk
212     if kk == 0:
213         inci_bot[ek,0] = 2*(k**2) + 4*k + 1
214         inci_bot[ek,1] = 6*k*(k+1) + 4*Ns - M - Mlb
215         inci_bot[ek,2] = 6*k*(k+1) + 4*Ns - 1 - M - Mlb
216         inci_bot[ek,3] = 2*(k**2)
217     else:
218         inci_bot[ek,0] = 6*k*(k+1) + 2*Ns + 1 - M - Mlb
219         inci_bot[ek,1] = 6*k*(k+1) + 2*Ns - M - Mlb
220         inci_bot[ek,2] = 2*(k**2) + 4*k + 1 - Mlb
221         inci_bot[ek,3] = 2*(k**2) + 2*k
222 for k in range(1,Ns-1):
223     kkk = np.tile(np.arange(4*k),(2,1))
224     kkk[0,k:] = kkk[0,k:] + 2
225     kkk[0,3*k:] = kkk[0,3*k:] + 2
226     for kk in range(3):
227         ek = 2*(k**2) + kk + k*((kk*(kk+1))/2)
228         etype_bot[ek:ek+((2-kk)*kk+1)*k] = 9 + kk
229         n0 = 1 + 5*k + 2*(k-1)*k
230         n1 = n0 + 1
231         n2 = 2 + 5*(k-1) + 2*(k-2)*(k-1)
232         n3 = n2 - 1
233         inci_bot[ek:ek+((2-kk)*kk+1)*k,0] = n0 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
234         inci_bot[ek:ek+((2-kk)*kk+1)*k,1] = n1 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
235         inci_bot[ek:ek+((2-kk)*kk+1)*k,2] = n2 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
236         inci_bot[ek:ek+((2-kk)*kk+1)*k,3] = n3 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
237 k = Ns-1
238 kkk = np.tile(np.arange(4*k),(2,1))
239 kkk[0,k:3*k] = np.arange(0,-2*Ns+2,-1)
240 kkk[0,3*k:] = np.arange(k,0,-1) - Mlb
241 for kk in range(3):
242     ek = 2*(k**2) + kk + k*((kk*(kk+1))/2)
243     etype_bot[ek:ek+((2-kk)*kk+1)*k] = 9 + kk
244     if kk == 1:
245         n0 = 4*Ns - 1 + 6*k*(k+1) - M - Mlb
246         n1 = n0 - 1
247     else:
248         n0 = 1 + 5*k + 2*(k-1)*k
249         n1 = n0 + 1 - kk
250         n2 = 2 + 5*(k-1) + 2*(k-2)*(k-1)

```

```

251     n3 = n2 - 1
252     inci_bot[ek:ek+((2-kk)*kk+1)*k,0] = n0 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
253     inci_bot[ek:ek+((2-kk)*kk+1)*k,1] = n1 + kkk[0,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
254     inci_bot[ek:ek+((2-kk)*kk+1)*k,2] = n2 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
255     inci_bot[ek:ek+((2-kk)*kk+1)*k,3] = n3 + kkk[1,((kk*(kk+1))/2)*k:((-kk**2)+5*kk+2)//2)*k]
256     inci_bot = inci_bot + M + Mlb
257     sym_bot = np.ndarray((N,2),dtype=np.uint32)
258     for k in range(Ns):
259         for kk in range(2):
260             ek = 2*((k+kk)**2) + k - kk*(2*k+1)
261             sym_bot[N-1-k*(Ns+1):N-k*Ns,kk] = np.arange(ek,ek+(1-2*kk)*(k+1),1-2*kk)
262     for k in range(Ns-1):
263         for kk in range(2):
264             sym_bot[(k+1)*Ns+k:Ns,kk] = np.arange(sym_bot[k*Ns+k,kk]+2*kk-1,sym_bot[k*Ns+k,kk]+(2*kk-1)*(Ns-k),2*kk-1)
265     sym_bot = sym_bot + 8*N
266     return coor_lb, coor_bot, inci_lb, inci_bot, sym_lb, sym_bot

```

3.3.4 ./source/python/elem.py

This script generates the local elemental stiffness matrices for each type of quadrilateral element, and for each type of augmented element. Firstly, the numpy module is imported.

```

1 import numpy as np

```

The function “getJ” is defined. Considering an isoparametric bilinear quadrilateral element, whose coordinates are stored in the matrix “sv”, for given coordinates “xi” and “eta” (defined in $[-1, 1] \times [-1, 1]$), the Jacobian matrix, its inverse and its determinant are computed.

```

2 def getJ(xi,eta,sv):
3     dN = 0.25*np.array([[(-1-eta), (1-eta), (1+eta), -(1+eta)],
4                          [-(1-xi), -(1+xi), (1+xi), (1-xi)]])
5     J = dN @ sv
6     Jdet = J[0,0]*J[1,1] - J[0,1]*J[1,0]
7     Jinv = (1/Jdet) * np.array([[ J[1,1], -J[0,1]],
8                                 [-J[1,0], J[0,0]]])
9     return Jinv, Jdet

```

The function “getB” is defined. Considering the quadrilateral element defined by the nodal coordinates “sv”, for given coordinates “xi” and “eta”, the matrix “Bmat” is computed. When it is applied on the nodal displacements vector, it yields the strain vector (in Voigt notation) at the position defined by “xi” and “eta”.

```

10 def getB(xi,eta,sv):
11     Jinv, Jdet = getJ(xi,eta,sv)
12     Ai = np.array([[1,0,0,0],
13                   [0,0,0,1],
14                   [0,1,1,0]])
15     Aj = np.zeros((4,4))
16     Aj[0:2,0:2] = Jinv
17     Aj[2:4,2:4] = Jinv
18     Ak = 0.25*np.array([[(-1-eta), 0.0, (1-eta), 0.0, (1+eta), 0.0, -(1+eta), 0.0],
19                          [-(1-xi), 0.0, -(1+xi), 0.0, (1+xi), 0.0, (1-xi), 0.0],
20                          [0.0, -(1-eta), 0.0, (1-eta), 0.0, (1+eta), 0.0, -(1+eta)],
21                          [0.0, -(1-xi), 0.0, -(1+xi), 0.0, (1+xi), 0.0, (1-xi)]])
22     Bmat = Ai @ Aj @ Ak
23     return Bmat, Jdet

```

The function “get_emat” is defined. Given the Young’s modulus (“Ey”) and the Poisson’s ratio (“nu”) of an isotropic material in plane stress state, the corresponding constitutive matrix is stored in “Ce”. Since the elemental stiffness matrix does not depend on the scale of the element, elements with shorter sides of 1.0 m and longer sides of $\sqrt{3}m$ are considered. The stiffness matrices, for each of the twelve types of quadrilateral elements, are computed through Gaussian quadrature with 2×2 points. They are stored in the tensor “Ket”, of dimensions $12 \times 8 \times 8$.

```

24 def get_emat(Ey,nu):
25     # constitutive matrix
26     Ce = (Ey/(1-nu**2))*np.array([[1,nu,0],[nu,1,0],[0,0,(1-nu)/2]])
27     # elements nodes
28     sv = np.ndarray((12,4,2))
29     s3 = np.sqrt(3)
30     sv[0,[:,,:]] = np.array([[ -0.5, 0.5*s3],
31                               [ 0.0, 0.0 ]],

```

```

32         [ 1.0, 0.0 ],
33         [ 1.0, s3]])
34 sv[1, :, :] = np.array([[ -1.0, 0.0 ],
35                          [ 0.0, 0.0 ],
36                          [ 0.5, 0.5*s3],
37                          [ -1.0, s3]])
38 sv[2, :, :] = np.array([[ -0.5, -0.5*s3],
39                          [ 0.0, 0.0 ],
40                          [ -0.5, 0.5*s3],
41                          [ -2.0, 0.0 ]])
42 sv[3, :, :] = np.array([[ 0.5, -0.5*s3],
43                          [ 0.0, 0.0 ],
44                          [ -1.0, 0.0 ],
45                          [ -1.0, -s3]])
46 sv[4, :, :] = np.array([[ 1.0, 0.0 ],
47                          [ 0.0, 0.0 ],
48                          [ -0.5, -0.5*s3],
49                          [ 1.0, -s3]])
50 sv[5, :, :] = np.array([[ 0.5, 0.5*s3],
51                          [ 0.0, 0.0 ],
52                          [ 0.5, -0.5*s3],
53                          [ 2.0, 0.0 ]])
54 sv[6, :, :] = np.array([[ 0.0, 0.0 ],
55                          [ 0.5, -0.5*s3],
56                          [ 2.0, 0.0 ],
57                          [ 1.5, 0.5*s3]])
58 sv[7, :, :] = np.array([[ 0.0, 0.0 ],
59                          [ 1.0, 0.0 ],
60                          [ 1.0, s3],
61                          [ 0.0, s3]])
62 sv[8, :, :] = np.array([[ 0.0, 0.0 ],
63                          [ 0.5, 0.5*s3],
64                          [ -1.0, s3],
65                          [ -1.5, 0.5*s3]])
66 sv[9, :, :] = np.array([[ 0.0, 0.0 ],
67                          [ -0.5, 0.5*s3],
68                          [ -2.0, 0.0 ],
69                          [ -1.5, -0.5*s3]])
70 sv[10, :, :] = np.array([[ 0.0, 0.0 ],
71                          [ -1.0, 0.0 ],
72                          [ -1.0, -s3],
73                          [ 0.0, -s3]])
74 sv[11, :, :] = np.array([[ 0.0, 0.0 ],
75                          [ -0.5, -0.5*s3],
76                          [ 1.0, -s3],
77                          [ 1.5, -0.5*s3]])
78 # stiffness matrices
79 Ket = np.ndarray((12,8,8))
80 gpoint = s3/3
81 for ek in range(12):
82     B0, J0 = getB(-gpoint, -gpoint, sv[ek, :, :])
83     B1, J1 = getB(gpoint, -gpoint, sv[ek, :, :])
84     B2, J2 = getB(gpoint, gpoint, sv[ek, :, :])
85     B3, J3 = getB(-gpoint, gpoint, sv[ek, :, :])
86     Ket[ek, :, :] = J0 * B0.T @ Ce @ B0 + J1 * B1.T @ Ce @ B1 + J2 * B2.T @ Ce @ B2 + J3 * B3.T @ Ce @ B3
87     Ket[ek, :, :] = 0.5*(Ket[ek, :, :]+Ket[ek, :, :].T)
88 return Ket

```

The function “get_augmat” is defined. It computes the stiffness variation matrices, stored in “dKelist”, and their factorizations (Equation 27), stored in “Hlist”, for the six different types of augmented elements.

```

89 def get_augmat(Ns, inci, etype, sym, dKe, small=1e-14):
90     N = Ns**2 # number of elements in the design domain
91     # different unconstrained augmented elements
92     aug_etype = np.ndarray((N), dtype=np.uint8)
93     aug_etype[0:N-1:Ns+1] = 0
94     for k in range(Ns-2):
95         aug_etype[k*(Ns+1)+1:(k+1)*Ns-1] = 1
96         aug_etype[(k+1)*(Ns+1)-1:N-Ns-1:Ns] = 2
97     aug_etype[Ns-1:N-1:Ns] = 3
98     aug_etype[N-Ns:N-1] = 4
99     aug_etype[N-1] = 5
100     # factorization
101     Hlist = []
102     dKelist = []
103     eledofs = np.ndarray((6,8), dtype=np.uint32)
104     locdofs = np.ndarray((6,8), dtype=np.uint32)
105     for e in [0,1,Ns,Ns-1,N-Ns,N-1]:
106         for k in range(6):
107             et = sym[e,k]
108             n0 = inci[et,0]
109             n1 = inci[et,1]
110             n2 = inci[et,2]
111             n3 = inci[et,3]
112             eledofs[k, :] = np.array([2*n0, 2*n0+1, 2*n1, 2*n1+1, 2*n2, 2*n2+1, 2*n3, 2*n3+1])
113             size = len(np.unique(eledofs))
114             elebool = np.ones((6,8), dtype=bool)

```



```

115     kkk = 0
116     for k in range(6):
117         for kk in range(8):
118             if elebool[k, kk]:
119                 mask = (eledofs==eledofs[k, kk])
120                 locdofs[mask] = kkk
121                 elebool[mask] = False
122                 kkk = kkk + 1
123     aug_dKe = np.zeros((size, size))
124     for k in range(6):
125         et = sym[e, k]
126         ek = etype[et]
127         grid = np.ix_(locdofs[k, :], locdofs[k, :])
128         aug_dKe[grid] = aug_dKe[grid] + dKe[ek, :, :]
129     D, V = np.linalg.eigh(aug_dKe)
130     mask = abs(D) > small
131     D = D[mask]
132     V = V[:, mask]
133     He = V*np.sqrt(D)
134     Hlist = Hlist + [He]
135     dKelist = dKelist + [aug_dKe]
136     return aug_etype, Hlist, dKelist

```

3.3.5 ./source/python/filters.py

This script generates the matrices used in the filtering procedures: smoothing sensitivity maps; and applying morphological operators on the density map. Firstly, the necessary modules are imported.

```

1  import numpy as np
2  from scipy.sparse import coo_matrix

```

The function “get_sfil” is defined. According to the conical filter radius (“rsen”), the matrix “Sf” is computed. The filtered sensitivity vectors are obtained by applying this matrix on the raw sensitivity vectors.

```

3  def get_sfil(N, sym, elepos, Q, rsen):
4      clist = []
5      csize = np.ndarray((N), dtype=np.uint32)
6      for e in range(N):
7          et = sym[e, 0]
8          c = np.argwhere(np.sum((elepos[et, :] - elepos)**2, axis=1) <= rsen**2)
9          clist = clist + [c[:, 0]]
10         csize[e] = len(c)
11     size = sum(csize)
12     row = np.ndarray((size), dtype=np.uint32)
13     col = np.ndarray((size), dtype=np.uint32)
14     data = np.ndarray((size))
15     i = 0
16     for e in range(N):
17         et = sym[e, 0]
18         c = clist[e]
19         num = csize[e]
20         weights = rsen - np.linalg.norm(elepos[et, :] - elepos[c, :], axis=1)
21         weights = weights/sum(weights)
22         row[i:i+num] = np.repeat(e, num)
23         col[i:i+num] = c
24         data[i:i+num] = weights
25         i = i + num
26     Sf = coo_matrix((data, (row, col)), shape=(N, 10*N))
27     Sf = Sf.tocsr()
28     Sf = Sf @ Q
29     return Sf

```

The function “get_mope” is defined. According to the radius “rmor”, the matrix “Mf” is computed. It is used to identify elements within range, when applying the morphological operators (erosion or dilation).

```

30 def get_mope(N, sym, elepos, Q, rmor):
31     clist = []
32     csize = np.ndarray((N), dtype=np.uint32)
33     for e in range(N):
34         et = sym[e, 0]
35         c = np.argwhere(np.sum((elepos[et, :] - elepos)**2, axis=1) <= rmor**2)
36         clist = clist + [c[:, 0]]
37         csize[e] = len(c)
38     size = sum(csize)
39     row = np.ndarray((size), dtype=np.uint32)
40     col = np.ndarray((size), dtype=np.uint32)
41     data = np.ndarray((size))
42     i = 0

```

```

43     for e in range(N):
44         et = sym[e,0]
45         c = clist[e]
46         num = csize[e]
47         row[i:i+num] = np.repeat(e,num)
48         col[i:i+num] = c
49         data[i:i+num] = 1.0
50         i = i + num
51     Mf = coo_matrix((data,(row,col)),shape=(N,10*N))
52     Mf = Mf.tocsr()
53     Mf = Mf @ Q
54     return Mf

```

3.3.6 ./source/python/rem_islands.py

This script is used to identify disconnected structural components in the topology. Firstly, the necessary modules are imported.

```

1  import numpy as np
2  import sys

```

The recursion limit is increased, so that the recursive function is not interrupted before the procedure is concluded. The recursive function “visit” is defined. According to the neighbors matrix, it performs a depth-first search, visiting all solids which are connected to a given reference element. After the search is concluded, the boolean mask “continent” identifies all solids that are connected to the reference element. While solids that are not in “continent” correspond to structural components disconnected from the reference element.

```

3  sys.setrecursionlimit(100000)
4  def visit(e,x,continent,neighbors):
5      continent[e] = True
6      for ee in neighbors[e,:]:
7          if x[ee] and (not continent[ee]):
8              visit(ee,x,continent,neighbors)
9      return

```

The function “get_neighbors” is defined. It computes the neighbors matrix. For each design variable, it stores the indices of the four design variables that correspond to directly connected neighbors. For elements on the boundaries of the design domain, some of these indices may be repeated. This does not hamper the considered procedure, since repeated indices will be automatically ignored by the depth-first search algorithm.

```

10 def get_neighbors(Ns,inci,inci_lb,inci_bot,sym,sym_lb,sym_bot):
11     N = Ns**2
12     Mf = 1 + 6*Ns*(Ns+1) + Ns + 4*N
13     Nf = 10*N
14     finci = np.vstack((inci,inci_lb,inci_bot))
15     fsym = np.hstack((sym,sym_lb,sym_bot))
16     nodes = -np.ones((Mf,6),dtype=int)
17     counter = np.zeros(Mf,dtype=int)
18     for e in range(Nf):
19         for k in range(4):
20             n = finci[e,k]
21             nodes[n,counter[n]] = e
22             counter[n] += 1
23     neighbors_extended = -np.ones((Nf,4),dtype=int)
24     counter = np.zeros(Nf,dtype=int)
25     for e in range(Nf):
26         for k in range(4):
27             n = finci[e,k]
28             for kk in range(6):
29                 ee = nodes[n,kk]
30                 if (ee != e) and (ee != -1):
31                     if ee not in neighbors_extended[e,:]:
32                         if len(np.setdiff1d(finci[e,:],finci[ee,:],assume_unique=True)) == 2:
33                             neighbors_extended[e,counter[e]] = ee
34                             counter[e] += 1
35     neighbors_ext = neighbors_extended.copy()
36     for ef in range(Nf):
37         for k in range(4):
38             if neighbors_extended[ef,k] != -1:
39                 e = np.argwhere(fsym==neighbors_extended[ef,k])[0,0]
40                 neighbors_ext[ef,k] = e
41     neighbors = np.ndarray((N,4),dtype=int)
42     for k in range(N):
43         neighbors[k,:] = neighbors_ext[fsym[k,0],:]
44     return neighbors

```

3.3.7 ./source/python/topopt.py

This script defines basic operations required to perform topology optimizations: the update of the topology; and the computation of sensitivity values. Only the WS approach is considered in this script, the CGS approach is implemented in the cython script “silp_sens.pyx”, detailed in a following section. Firstly, the numpy module is imported.

```
1 import numpy as np
```

The function “update” is defined. According to the list of elements whose states must be switched (“elist”), the density vector (“x”) and the COO unconstrained global stiffness matrix (“Kg_coo”) are updated. If the flag “solve_sys” is *True*, the function returns the total displacements matrix (“Ug”) and the CSC constrained global stiffness matrix (“Kr”) of the updated topology. Otherwise, it returns nothing.

```
2 def update(x,etype,sym,pk,Ketvec,P,Kg_coo,Zr,shift,Uhat,factor,elist,solve_sys=True):
3     x[elist] = -x[elist]
4     for e in elist:
5         if x[e]:
6             for k in range(6):
7                 et = sym[e,k]
8                 ek = etype[et]
9                 Kg_coo.data[64*et:64*et+64] = Ketvec[ek,:]
10        else:
11            for k in range(6):
12                et = sym[e,k]
13                ek = etype[et]
14                Kg_coo.data[64*et:64*et+64] = pk*Ketvec[ek,:]
15    if solve_sys:
16        Kg_csc = Kg_coo.tocsc()
17        Kr = P.T @ Kg_csc @ P
18        Kr = Kr + shift*Zr
19        Kr.sort_indices()
20        Kr.data = Kr.data - shift*Zr.data
21        Fr = -P.T @ Kg_csc @ Uhat
22        factor.cholesky_inplace(Kr)
23        Ur = factor(Fr)
24        Ug = Uhat + P @ Ur
25        return Ug, Kr
26
```

The function “ws” is defined. It computes the exact sensitivity values of the diagonal terms of \mathbf{C} through WS approach.

```
27 def ws(x,aug_etype,sym,P,factor,inci,Ug,Hlist):
28     N = len(x)
29     dC00_w = np.ndarray((N))
30     dC11_w = np.ndarray((N))
31     dC22_w = np.ndarray((N))
32     eledofs = np.ndarray((6,8),dtype=np.uint32)
33     for e in range(N):
34         for k in range(6):
35             et = sym[e,k]
36             n0 = inci[et,0]
37             n1 = inci[et,1]
38             n2 = inci[et,2]
39             n3 = inci[et,3]
40             eledofs[k,:] = np.array([2*n0,2*n0+1,2*n1,2*n1+1,2*n2,2*n2+1,2*n3,2*n3+1])
41         aug_ek = aug_etype[e]
42         He = Hlist[aug_ek]
43         size = He.shape
44         elebool = np.ones((6,8),dtype=bool)
45         aug_eledofs = np.ndarray((size[0]),dtype=np.uint32)
46         kkk = 0
47         for k in range(6):
48             for kk in range(8):
49                 if elebool[k,kk]:
50                     mask = (eledofs==eledofs[k,kk])
51                     elebool[mask] = False
52                     aug_eledofs[kkk] = eledofs[k,kk]
53                     kkk = kkk + 1
54         Ie = np.eye(size[1])
55         Ue = Ug[aug_eledofs,:]
56         Pe = P[aug_eledofs,:]
57         Hr = Pe.T @ He
58         Ahalf = factor.solve_L(factor.apply_P(Hr),use_LDLt_decomposition=False)
59         Ae = Ahalf.T @ Ahalf
60         Ve = He.T @ Ue
61         if x[e]:
```

```

62         dCh = -Ve.T @ np.linalg.solve(Ie-Ae,Ve)
63     else:
64         dCh = Ve.T @ np.linalg.solve(Ie+Ae,Ve)
65     dC00_w[e] = dCh[0,0]
66     dC11_w[e] = dCh[1,1]
67     dC22_w[e] = dCh[2,2]
68     return dC00_w, dC11_w, dC22_w

```

3.3.8 ./source/python/ilp_solver.py

This script is used to solve each linearized subproblem. If there is no Young’s modulus constraint, the BESO algorithm is used, otherwise, the pulp module is used to solve the problem with branch-and-bound simplex. Firstly, the necessary modules are imported.

```

1  import numpy as np
2  import pulp as pp

```

The function “solve_ILP” is defined. It receives the number of design variables (“N”); the current topology vector (“x”); the sensitivity vector of the objective function (“alpha”); the sensitivity vector of the constraint function (“alpha_h”); the current value of the constraint function (“h_bar”); the tolerance of the constraint function (“h_lim”); the maximal topology variation (“dXmax”); the sense of the optimization (“sense”); and the sense of the constraint function (“sense_h”). Then, it returns the solution of the corresponding integer linear problem. Some optional parameter can be included: “tLim” sets a maximum time for the solver (in seconds); “rErr” sets a relative gap tolerance for the solver to stop; if “verbose” is *True* the solver’s log is shown.

```

3  def solve_ILP(N,x,alpha,alpha_h,h_bar,h_lim,dXmax,sense='min',sense_h='L',tLim=np.infty,rErr=None,verbose=False):
4      # Integer Linear Programming problem
5      if sense == 'min': # minimization problem
6          prob = pp.LpProblem('ILP', pp.LpMinimize)
7      else: # maximization problem
8          prob = pp.LpProblem('ILP', pp.LpMaximize)
9      variables = [f'{e:07d}' for e in range(N)]
10     xvars = pp.LpVariable.dicts('x',variables,cat='Binary')
11     # Objective Function
12     # f(x) = f(xbar) + alpha * [x-xbar]
13     # min/max { f(x) } = min/max { alpha * x }
14     alpha_obj = dict(zip(variables,alpha))
15     prob += pp.lpSum([alpha_obj[v] * xvars[v] for v in variables])
16     # General Constraint
17     # h(x) = h(xbar) + alpha_h * [x-xbar] <= (or >=) h_lim
18     # alpha_h * x <= (or >=) b_h
19     b_h = h_lim - h_bar + sum(alpha_h[x])
20     alpha_hcon = dict(zip(variables,alpha_h))
21     if sense_h == 'L': # constraint : <=
22         prob += pp.lpSum([alpha_hcon[v] * xvars[v] for v in variables]) <= b_h
23     else: # constraint : >=
24         prob += pp.lpSum([alpha_hcon[v] * xvars[v] for v in variables]) >= b_h
25     # Maximal Topological Change Constraint
26     # g(x) = ||x-xbar||_1 <= dXmax
27     # g(x) = g(xbar) + alpha_g * [x-xbar] <= dXmax
28     # alpha_g * x <= b_g
29     g_bar = 0.0
30     alpha_g = np.ones(N)
31     alpha_g[x] = -1.0
32     b_g = dXmax - g_bar + sum(alpha_g[x])
33     alpha_gcon = dict(zip(variables,alpha_g))
34     prob += pp.lpSum([alpha_gcon[v] * xvars[v] for v in variables]) <= b_g
35     # Solve ILP
36     prob.solve(solver=pp.COIN_CMD(msg=verbose, gapRel=rErr, timeLimit=tLim))
37     y = np.array([v.varValue for v in prob.variables()],dtype=bool)
38     if verbose:
39         print('linearized objective variation = {:.1e}'.format(sum(alpha[y])-sum(alpha[x])))
40     return y

```

The function “solve_BESO” is defined. It solves an integer linear problem of binary variables with a single constraint, given by the maximal topology variation (“dXmax”).

```

41  def solve_BESO(N,x,alpha,dXmax,sense='min'):
42      # Integer Linear Programming problem without extra constraints
43      y = x.copy()
44      if sense == 'min': # minimization problem
45          mask = (x & (alpha>0.0)) | (~x & (alpha<0.0))
46      else: # maximization problem
47          mask = (x & (alpha<0.0)) | (~x & (alpha>0.0))
48      Nv = sum(mask)

```

```

49     if Nv == 0:
50         return y
51     xsub = x[mask]
52     arg = np.argsort(abs(alpha[mask]))
53     if Nv < dXmax:
54         xsub = -xsub
55     else:
56         xsub[arg[-dXmax:]] = -xsub[arg[-dXmax:]]
57     y[mask] = xsub
58     return y

```

3.3.9 ./source/python/generate_metamat.py

This script concludes the generation of the dataset. It verifies if the generated data is coherent, that is, there is no redundant data and all expected files exist. Then, the data is transferred to the “dataset” folder. Output subfolders are renamed so that a unique number is attributed to each one of them. Duplicate files are deleted, but all folders and logs are preserved.

Firstly, the necessary modules are imported. The number of optimizations stored in the same file (“noptf”) is set to the same value used when generating the data, which is 7.

```

1  import os, sys, shutil
2  noptf = 7 # number of optimizations stored in the same file

```

The data generated by the program “basecell_silp” is verified. The script checks if: the output folder exists; there is at least one subfolder in it; no duplicated data have been generated; logs subfolders exist; the number of performed optimizations is a multiple of the “noptf” parameter (although this is not necessary, since the generation was performed in groups of exactly “noptf” cases, this is verified to make sure that everything is in order); the number of generated files is coherent; exactly 27 arrays of data have been written in each files subfolder.

```

3  if not os.path.exists('./SILP/output'):
4      print('no output directory')
5      sys.exit()
6  runs = sorted(os.listdir('./SILP/output'))
7  if len(runs) == 0:
8      print('no runs have been found')
9      sys.exit()
10 total = 0
11 r2 = -1
12 for k in range(len(runs)):
13     run_dir = './SILP/output/' + runs[k] + '/'
14     r1 = int(runs[k][4:9])
15     if r1 <= r2:
16         print('redundant runs : ' + runs[k])
17         sys.exit()
18     r2 = int(runs[k][10:])
19     rnum = r2 - r1 + 1
20     files = sorted(os.listdir(run_dir))
21     if files[-1] == 'logs':
22         files = files[:-1]
23     else:
24         print('missing logs directory : ' + runs[k])
25         sys.exit()
26     if rnum % noptf != 0:
27         print('number of cases is not a multiple of noptf : ' + runs[k])
28         sys.exit()
29     if noptf*len(files) != rnum:
30         print('incoherent number of files : ' + runs[k])
31         sys.exit()
32     total += rnum
33     for kk in range(len(files)):
34         file_dir = run_dir + files[kk]
35         if len(os.listdir(file_dir)) != 27:
36             print('wrong number of files : ' + runs[k] + '/' + files[kk])
37             sys.exit()
38 if total % noptf != 0:
39     print('something is wrong...')
40     sys.exit()
41 total = total // noptf
42 print('valid SILP dataset!')
43 print('{:04d} / {:04d} files in the SILP dataset ({:5.1f} %)\n'.format(
44     total, 18382//noptf, total*noptf*100/18382))

```

Then, the script verifies if there is already a previously generated dataset in the “dataset” folder. In order to avoid undesired overwriting, it is expected that folders be renamed when generating multiple datasets.

```

45 # check directories
46 if not os.path.exists('.././dataset'):
47     os.mkdir('.././dataset')
48 if not os.path.exists('.././dataset/SILP'):
49     os.mkdir('.././dataset/SILP')
50 else:
51     if len(os.listdir('.././dataset/SILP')) > 0:
52         print('a SILP dataset has already been generated, rename its directory before generating a new one')
53         sys.exit()

```

If everything is in order, the data is transferred to the “dataset” folder. All generated data is transferred, subfolders are renamed according to the “global_id” variable, so that a unique number is attributed to each files subfolder.

```

54 global_id = 0
55 runs = sorted(os.listdir('../SILP/output'))
56 for k in range(len(runs)):
57     run_dir = '../SILP/output/' + runs[k] + '/'
58     files = sorted(os.listdir(run_dir))
59     files = files[:-1]
60     for kk in range(len(files)):
61         if global_id % 100 == 0:
62             print('{:04d} / {:04d} : files have been moved to the SILP dataset ({:5.1f} %)' .format(
63                 global_id, total, global_id*100/total))
64             file_dir = run_dir + files[kk] + '/'
65             destination = '.././dataset/SILP/f{:04d}' .format(global_id)
66             os.mkdir(destination)
67             for file in os.listdir(file_dir):
68                 source = file_dir + file
69                 shutil.move(source, destination)
70             global_id += 1
71 print('{:04d} / {:04d} : files have been moved to the SILP dataset ({:5.1f} %)' .format(
72     global_id, total, global_id*100/total))
73 print('[ SILP dataset generated ]')

```

3.4 Implementation – Cython

3.4.1 ./source/cython/cython_setup.py

This python script compiles the cython script: “silp_sens.pyx”.

```

1 from setuptools import setup
2 from Cython.Build import cythonize
3 setup(
4     ext_modules = cythonize(
5         ['./silp_sens.pyx'],
6         compiler_directives={'language_level' : "3"},
7         annotate=False)
8 )

```

3.4.2 ./source/cython/silp_sens.pyx

This script performs the CGS analysis for the diagonal terms of the elasticity matrix of the homogenized metamaterial (\mathbf{C}). Firstly, some flags are set and the cython module is imported.

```

1 # cython: boundscheck=False
2 # cython: wraparound=False
3 # cython: cdivision=True
4 cimport cython

```

The function “cython_cgs” is defined. It performs the CGS-0, CGS-1 and CGS-2 sensitivity analyses for the diagonal terms of \mathbf{C} . The function returns nothing, the sensitivity vectors “dC00_0”, “dC11_0”, “dC22_0”, “dC00_1”, “dC11_1”, “dC22_1”, “dC00_2”, “dC11_2” and “dC22_2” are received as input then edited during execution (the changes are preserved in the outer scope). The approximations are computed in sequence: CGS-0 first (“dC00_0”, “dC11_0” and “dC22_0”); then CGS-1 (“dC00_1”, “dC11_1” and “dC22_1”); and lastly CGS-2 (“dC00_2”, “dC11_2” and “dC22_2”).

The matrix “P” is used to constrain the elemental matrices. For each augmented element, the global matrix is sliced and the resulting submatrix is stored in the variable “Kbb”, which can have dimensions up to 192×192 .

The CGS-0 sensitivity values are easily obtained from the elemental strain energy values. The CGS-1 and CGS-2 sensitivity values are obtained after computing the required coefficients: ω_{hm} (“whm”), ω_{mk} (“wmk”), ϕ_{m1} (“pm1”), ϕ_{m2} (“pm2”) and $\phi_{\eta 2}$ (“peta2”).

```

5  cdef void cython_cgs(double[:] dC00_0, double[:] dC11_0, double[:] dC22_0, double[:] dC00_1, double[:] dC11_1,
6      double[:] dC22_1, double[:] dC00_2, double[:] dC11_2, double[:] dC22_2, long long[:] x, long long N,
7      long long[:,::1] sym, long long[:] etype, long long[:] aug_etype, long long[:,::1] inci, double[:,::1] Ug,
8      double[:,::1] dKe, long long[:] Pindices, long long[:] Pindptr, long long[:] Kindices,
9      long long[:] Kindptr, double[:] Kdata, double[:,::1] aug_dKe0, double[:,::1] aug_dKe1,
10     double[:,::1] aug_dKe2, double[:,::1] aug_dKe3, double[:,::1] aug_dKe4, double[:,::1] aug_dKe5):
11     cdef long long e
12     cdef long long i
13     cdef long long j
14     cdef long long k
15     cdef long long kk
16     cdef long long kkk
17     cdef long long et
18     cdef long long ek
19     cdef long long aug_ek
20     cdef long long n0
21     cdef long long n1
22     cdef long long n2
23     cdef long long n3
24     cdef long long dof
25     cdef long long ptr0
26     cdef long long ptr1
27     cdef long long pt0
28     cdef long long pt1
29     cdef long long ptm
30     cdef long long bvar
31     cdef long long row
32     cdef long long col
33     cdef long long dsize
34     cdef long long nPids
35     cdef long long size_h
36     cdef long long nKids
37     cdef long long size_k
38     cdef long long verylarge
39     cdef long long aug_eledofs[48]
40     cdef long long Peindptr[49]
41     cdef long long Peindices[48]
42     cdef long long Pe_h[48]
43     cdef long long dofs_h[48]
44     cdef long long uniquebool[864]
45     cdef long long Kreindices[864]
46     cdef long long dofs_k[192]
47     cdef long long dofs_kk[192]
48     cdef long long elebool[6][8]
49     cdef long long eledofs[6][8]
50     cdef double dCdx[3]
51     cdef double divisor[3]
52     cdef double pm1[3]
53     cdef double pm2[3]
54     cdef double peta2[3]
55     cdef double whm[3]
56     cdef double wmk[3]
57     cdef double wketa[3]
58     cdef double wetaxi[3]
59     cdef double Minv[192]
60     cdef double Kbb[192][192]
61     cdef double auxmat[48][48]
62     cdef double Ue[48][3]
63     cdef double zh[48][3]
64     cdef double zm[48][3]
65     cdef double zk[192][3]
66     cdef double zeta[192][3]
67     cdef double zxi[192][3]
68     verylarge = 9223372036854775807
69     for e in range(N):
70         for k in range(3):
71             dCdx[k] = 0.0
72         for k in range(6):
73             et = sym[e][k]
74             ek = etype[et]
75             n0 = inci[et][0]
76             n1 = inci[et][1]
77             n2 = inci[et][2]
78             n3 = inci[et][3]
79             eledofs[k][0] = 2*n0
80             eledofs[k][1] = 2*n0+1
81             eledofs[k][2] = 2*n1
82             eledofs[k][3] = 2*n1+1
83             eledofs[k][4] = 2*n2
84             eledofs[k][5] = 2*n2+1
85             eledofs[k][6] = 2*n3
86             eledofs[k][7] = 2*n3+1
87         for kk in range(8):
88             Ue[kk][0] = Ug[eledofs[k][kk]][0]

```

```

89         Ue[kk][1] = Ug[eledofs[k][kk]][1]
90         Ue[kk][2] = Ug[eledofs[k][kk]][2]
91     for i in range(8):
92         for j in range(8):
93             dCdx[0] = dCdx[0] + Ue[i][0]*dKe[ek][i][j]*Ue[j][0]
94             dCdx[1] = dCdx[1] + Ue[i][1]*dKe[ek][i][j]*Ue[j][1]
95             dCdx[2] = dCdx[2] + Ue[i][2]*dKe[ek][i][j]*Ue[j][2]
96 if x[e] == 1:
97     dC00_0[e] = -dCdx[0]
98     dC11_0[e] = -dCdx[1]
99     dC22_0[e] = -dCdx[2]
100 else:
101     dC00_0[e] = dCdx[0]
102     dC11_0[e] = dCdx[1]
103     dC22_0[e] = dCdx[2]
104 aug_ek = aug_etype[e]
105 if aug_ek < 3:
106     dsize = 48
107 elif aug_ek < 5:
108     dsize = 36
109 else:
110     dsize = 26
111 for k in range(6):
112     for kk in range(8):
113         elebool[k][kk] = 1
114 kkk = 0
115 for k in range(6):
116     for kk in range(8):
117         if elebool[k][kk] == 1:
118             for i in range(6):
119                 for j in range(8):
120                     if eledofs[i][j] == eledofs[k][kk]:
121                         elebool[i][j] = 0
122                         aug_eledofs[kkk] = eledofs[k][kk]
123                         kkk = kkk + 1
124 nPids = 0
125 Peindptr[0] = 0
126 for k in range(dsize):
127     Ue[k][0] = Ug[aug_eledofs[k]][0]
128     Ue[k][1] = Ug[aug_eledofs[k]][1]
129     Ue[k][2] = Ug[aug_eledofs[k]][2]
130     ptr0 = Pindptr[aug_eledofs[k]]
131     ptr1 = Pindptr[aug_eledofs[k]+1]
132     if ptr1 > ptr0:
133         Peindptr[k+1] = Peindptr[k] + 1
134         Peindices[nPids] = Pindices[ptr0]
135         nPids = nPids + 1
136     else:
137         Peindptr[k+1] = Peindptr[k]
138 for k in range(nPids):
139     uniquebool[k] = 1
140 size_h = 0
141 for k in range(nPids):
142     if uniquebool[k] == 1:
143         for kk in range(nPids):
144             if Peindices[kk] == Peindices[k]:
145                 uniquebool[kk] = 0
146                 dofs_h[size_h] = Peindices[k]
147                 size_h = size_h + 1
148 nKids = 0
149 for k in range(size_h):
150     ptr0 = Kindptr[dofs_h[k]]
151     ptr1 = Kindptr[dofs_h[k]+1]
152     for kk in range(ptr1-ptr0):
153         Kreindices[nKids+kk] = Kindices[ptr0+kk]
154     nKids = nKids + (ptr1-ptr0)
155 for k in range(nKids):
156     uniquebool[k] = 1
157 size_k = 0
158 for k in range(nKids):
159     if uniquebool[k] == 1:
160         for kk in range(k, nKids):
161             if Kreindices[kk] == Kreindices[k]:
162                 uniquebool[kk] = 0
163                 dofs_k[size_k] = Kreindices[k]
164                 size_k = size_k + 1
165 for k in range(size_h):
166     dofs_kk[k] = dofs_h[k]
167 kkk = size_h
168 for k in range(size_k):
169     bvar = 1
170     for kk in range(size_h):
171         if dofs_k[k] == dofs_h[kk]:
172             bvar = 0
173             break
174     if bvar == 1:
175         dofs_kk[kkk] = dofs_k[k]
176         kkk = kkk + 1
177 for k in range(dsize):
178     Pe_h[k] = verylarge
179 row = 0

```



```

180     for k in range(size_h):
181         for kk in range(nPids):
182             if Peindices[kk] == dofs_h[k]:
183                 for kkk in range(dsize):
184                     if kk < Peindptr[kkk+1]:
185                         row = kkk
186                         break
187                 Pe_h[row] = k
188     for i in range(size_h):
189         for k in range(dsize):
190             auxmat[i][k] = 0.0
191     if aug_ek == 0:
192         for j in range(dsize):
193             i = Pe_h[j]
194             if i != verylarge:
195                 for k in range(dsize):
196                     auxmat[i][k] = auxmat[i][k] + aug_dKe0[j][k]
197     elif aug_ek == 1:
198         for j in range(dsize):
199             i = Pe_h[j]
200             if i != verylarge:
201                 for k in range(dsize):
202                     auxmat[i][k] = auxmat[i][k] + aug_dKe1[j][k]
203     elif aug_ek == 2:
204         for j in range(dsize):
205             i = Pe_h[j]
206             if i != verylarge:
207                 for k in range(dsize):
208                     auxmat[i][k] = auxmat[i][k] + aug_dKe2[j][k]
209     elif aug_ek==3:
210         for j in range(dsize):
211             i = Pe_h[j]
212             if i != verylarge:
213                 for k in range(dsize):
214                     auxmat[i][k] = auxmat[i][k] + aug_dKe3[j][k]
215     elif aug_ek==4:
216         for j in range(dsize):
217             i = Pe_h[j]
218             if i != verylarge:
219                 for k in range(dsize):
220                     auxmat[i][k] = auxmat[i][k] + aug_dKe4[j][k]
221     else:
222         for j in range(dsize):
223             i = Pe_h[j]
224             if i != verylarge:
225                 for k in range(dsize):
226                     auxmat[i][k] = auxmat[i][k] + aug_dKe5[j][k]
227     for i in range(size_h):
228         zh[i][0] = 0.0
229         zh[i][1] = 0.0
230         zh[i][2] = 0.0
231         for j in range(dsize):
232             zh[i][0] = zh[i][0] + auxmat[i][j] * Ue[j][0]
233             zh[i][1] = zh[i][1] + auxmat[i][j] * Ue[j][1]
234             zh[i][2] = zh[i][2] + auxmat[i][j] * Ue[j][2]
235     for col in range(size_k):
236         ptr0 = Kindptr[dofs_kk[col]]
237         ptr1 = Kindptr[dofs_kk[col]+1]
238         for row in range(col,size_k):
239             dof = dofs_kk[row]
240             if (dof < Kindices[ptr0]) or (dof > Kindices[ptr1-1]):
241                 Kbb[row][col] = 0.0
242             else:
243                 pt0 = ptr0
244                 pt1 = ptr1
245                 bvar = 1
246                 while (pt1-pt0) > 1:
247                     ptm = (pt0+pt1)//2
248                     if dof > Kindices[ptm]:
249                         pt0 = ptm
250                     elif dof < Kindices[ptm]:
251                         pt1 = ptm
252                     else:
253                         Kbb[row][col] = Kdata[ptm]
254                         bvar = 0
255                         break
256                 if bvar == 1:
257                     if dof == Kindices[pt0]:
258                         Kbb[row][col] = Kdata[pt0]
259                     else:
260                         Kbb[row][col] = 0.0
261     if x[e] == 1:
262         for j in range(dsize):
263             k = Pe_h[j]
264             for i in range(size_h):
265                 if k <= i:
266                     Kbb[i][k] = Kbb[i][k] - auxmat[i][j]
267     else:
268         for j in range(dsize):
269             k = Pe_h[j]
270             for i in range(size_h):

```

```

271         if k <= i:
272             Kbb[i][k] = Kbb[i][k] + auxmat[i][j]
273     for k in range(size_k):
274         Minv[k] = 1.0/Kbb[k][k]
275     for k in range(size_h):
276         zm[k][0] = Minv[k] * zh[k][0]
277         zm[k][1] = Minv[k] * zh[k][1]
278         zm[k][2] = Minv[k] * zh[k][2]
279         zk[k][0] = Kbb[k][k] * zm[k][0]
280         zk[k][1] = Kbb[k][k] * zm[k][1]
281         zk[k][2] = Kbb[k][k] * zm[k][2]
282     for i in range(size_h,size_k):
283         zk[i][0] = 0.0
284         zk[i][1] = 0.0
285         zk[i][2] = 0.0
286     for i in range(size_h):
287         for j in range(i):
288             zk[i][0] = zk[i][0] + Kbb[i][j] * zm[j][0]
289             zk[i][1] = zk[i][1] + Kbb[i][j] * zm[j][1]
290             zk[i][2] = zk[i][2] + Kbb[i][j] * zm[j][2]
291             zk[j][0] = zk[j][0] + Kbb[i][j] * zm[i][0]
292             zk[j][1] = zk[j][1] + Kbb[i][j] * zm[i][1]
293             zk[j][2] = zk[j][2] + Kbb[i][j] * zm[i][2]
294     for i in range(size_h,size_k):
295         for j in range(size_h):
296             zk[i][0] = zk[i][0] + Kbb[i][j] * zm[j][0]
297             zk[i][1] = zk[i][1] + Kbb[i][j] * zm[j][1]
298             zk[i][2] = zk[i][2] + Kbb[i][j] * zm[j][2]
299     for i in range(size_k):
300         zeta[i][0] = Minv[i] * zk[i][0]
301         zeta[i][1] = Minv[i] * zk[i][1]
302         zeta[i][2] = Minv[i] * zk[i][2]
303         zxi[i][0] = Kbb[i][i] * zeta[i][0]
304         zxi[i][1] = Kbb[i][i] * zeta[i][1]
305         zxi[i][2] = Kbb[i][i] * zeta[i][2]
306     for i in range(size_k):
307         for j in range(i):
308             zxi[i][0] = zxi[i][0] + Kbb[i][j] * zeta[j][0]
309             zxi[i][1] = zxi[i][1] + Kbb[i][j] * zeta[j][1]
310             zxi[i][2] = zxi[i][2] + Kbb[i][j] * zeta[j][2]
311             zxi[j][0] = zxi[j][0] + Kbb[i][j] * zeta[i][0]
312             zxi[j][1] = zxi[j][1] + Kbb[i][j] * zeta[i][1]
313             zxi[j][2] = zxi[j][2] + Kbb[i][j] * zeta[i][2]
314     whm[0] = 0.0
315     whm[1] = 0.0
316     whm[2] = 0.0
317     wmk[0] = 0.0
318     wmk[1] = 0.0
319     wmk[2] = 0.0
320     for k in range(size_h):
321         whm[0] = whm[0] + zh[k][0]*zm[k][0]
322         whm[1] = whm[1] + zh[k][1]*zm[k][1]
323         whm[2] = whm[2] + zh[k][2]*zm[k][2]
324         wmk[0] = wmk[0] + zm[k][0]*zk[k][0]
325         wmk[1] = wmk[1] + zm[k][1]*zk[k][1]
326         wmk[2] = wmk[2] + zm[k][2]*zk[k][2]
327     pm1[0] = whm[0]/wmk[0]
328     pm1[1] = whm[1]/wmk[1]
329     pm1[2] = whm[2]/wmk[2]
330     if x[e] == 1:
331         dC00_1[e] = -dCdx[0] - pm1[0]*whm[0]
332         dC11_1[e] = -dCdx[1] - pm1[1]*whm[1]
333         dC22_1[e] = -dCdx[2] - pm1[2]*whm[2]
334     else:
335         dC00_1[e] = dCdx[0] - pm1[0]*whm[0]
336         dC11_1[e] = dCdx[1] - pm1[1]*whm[1]
337         dC22_1[e] = dCdx[2] - pm1[2]*whm[2]
338     wketa[0] = 0.0
339     wketa[1] = 0.0
340     wketa[2] = 0.0
341     wetaxi[0] = 0.0
342     wetaxi[1] = 0.0
343     wetaxi[2] = 0.0
344     for k in range(size_k):
345         wketa[0] = wketa[0] + zk[k][0]*zeta[k][0]
346         wketa[1] = wketa[1] + zk[k][1]*zeta[k][1]
347         wketa[2] = wketa[2] + zk[k][2]*zeta[k][2]
348         wetaxi[0] = wetaxi[0] + zeta[k][0]*zxi[k][0]
349         wetaxi[1] = wetaxi[1] + zeta[k][1]*zxi[k][1]
350         wetaxi[2] = wetaxi[2] + zeta[k][2]*zxi[k][2]
351     divisor[0] = wmk[0]*wetaxi[0]-wketa[0]*wketa[0]
352     divisor[1] = wmk[1]*wetaxi[1]-wketa[1]*wketa[1]
353     divisor[2] = wmk[2]*wetaxi[2]-wketa[2]*wketa[2]
354     pm2[0] = (whm[0]*wetaxi[0]-wmk[0]*wketa[0])/divisor[0]
355     pm2[1] = (whm[1]*wetaxi[1]-wmk[1]*wketa[1])/divisor[1]
356     pm2[2] = (whm[2]*wetaxi[2]-wmk[2]*wketa[2])/divisor[2]
357     peta2[0] = (wmk[0]*wmk[0]-whm[0]*wketa[0])/divisor[0]
358     peta2[1] = (wmk[1]*wmk[1]-whm[1]*wketa[1])/divisor[1]
359     peta2[2] = (wmk[2]*wmk[2]-whm[2]*wketa[2])/divisor[2]
360     if x[e] == 1:
361         dC00_2[e] = -dCdx[0] - (pm2[0]*whm[0]+peta2[0]*wmk[0])

```

```

362         dC11_2[e] = -dCdx[1] - (pm2[1]*whm[1]+peta2[1]*wmk[1])
363         dC22_2[e] = -dCdx[2] - (pm2[2]*whm[2]+peta2[2]*wmk[2])
364     else:
365         dC00_2[e] = dCdx[0] - (pm2[0]*whm[0]+peta2[0]*wmk[0])
366         dC11_2[e] = dCdx[1] - (pm2[1]*whm[1]+peta2[1]*wmk[1])
367         dC22_2[e] = dCdx[2] - (pm2[2]*whm[2]+peta2[2]*wmk[2])
368     return

```

The “cgs” function is defined to call the cython function in the python program. It receives as input: the vectors to which the sensitivity values will be assigned, “dC00_0”, “dC11_0”, “dC22_0”, “dC00_1”, “dC11_1”, “dC22_1”, “dC00_2”, “dC11_2” and “dC22_2”; the topology vector, “x”; the number of design variables, “N”; the symmetry matrix, “sym”; the vectors with element types, “etype” and “aug_etype”; the incidence matrix, “inci”; the unconstrained total displacements matrix, “Ug”; the elemental stiffness variation matrices, “dKe” and “dKelist”; the matrix used to apply the periodic boundary conditions, “P”; and the CSC constrained global stiffness matrix, “Kr”.

```

369 def cgs(dC00_0,dC11_0,dC22_0,dC00_1,dC11_1,dC22_1,dC00_2,dC11_2,dC22_2,x,N,sym,etype,aug_etype,inci,Ug,dKe,P,Kr,dKelist):
370     cython_cgs(dC00_0, dC11_0, dC22_0, dC00_1, dC11_1, dC22_1, dC00_2, dC11_2, dC22_2, x.astype("int64"),
371         N, sym.astype("int64"), etype.astype("int64"), aug_etype.astype("int64"),inci.astype("int64"), Ug, dKe,
372         P.indices.astype("int64"), P.indptr.astype("int64"), Kr.indices.astype("int64"), Kr.indptr.astype("int64"), Kr.data,
373         dKelist[0], dKelist[1], dKelist[2], dKelist[3], dKelist[4], dKelist[5])
374     return

```

3.5 Implementation – Sampling

3.5.1 ./sample/SILP/sample.py

This script generates figures for a selected sample of the dataset. For a given selection of files, png images for the topologies, sensitivity maps and displacements fields can be generated and saved in subfolders of the “sample/SILP” directory. Plots of the Poisson’s ratio, Young’s modulus and volume of material throughout a given optimization can also be generated and saved as png images.

Firstly, the necessary modules are imported.

```

1 import os, sys
2 import numpy as np
3 import matplotlib.pyplot as plt
4 import matplotlib.collections as clct
5 from adjust import adjust
6 sys.path.append('../source/python/SILP/')
7 from mesh import get_mesh

```

Then, the user selects which files should be read through the parameters “file_ini”, which specifies the index of the first file to be read, and “file_lim”, which specifies the file index limit (non inclusive). Boolean flags are used to specify which figures should be generated: if “fig_top_opt” is *True*, figures are generated for all optimized topologies; if “fig_top” is *True*, figures are generated for all topologies (not only the optimized); if “fig_sen” is *True*, figures are generated for all sensitivity vectors of the diagonal terms of **C**; if “fig_dis” is *True*, figures are generated for all displacements vectors; and if “fig_nu_Ey_vol” is *True*, figures are generated with all the plots of Poisson’s ratio, Young’s modulus and volume.

In order to avoid generating an unreasonable number of figures, users should be careful when defining these parameters. Each file contains the results of “noptf” optimizations, and each optimization produces several topology, sensitivity and displacements vectors (in average, 75 topology vectors, 900 sensitivity vectors and 225 displacements vectors).

```

8 file_ini      = 0      # initial file index |from file 0
9 file_lim      = 2      # file index limit   |up to file 2625
10 fig_top_opt   = True   # optimized topology
11 fig_top       = True   # topology vectors
12 fig_sen       = True   # sensitivity vectors
13 fig_dis       = True   # displacements vectors
14 fig_nu_Ey_vol = True   # Poisson's ratio, Young's modulus and volume

```

Some geometric properties are defined and the mesh is generated. The program checks if the “dataset” folder exists, then the folders in which the figures will be saved are created.

```

15 # fixed properties
16 Ns = 32 # number of elements in each side of the design domain
17 N = Ns**2 # number of elements in the design domain
18 Nt = 6*N # number of elements in the base cell
19 # area = 6*Lx*Ly = 1.0
20 Lx = 1.0/(108**0.25) # design domain shorter side
21 Ly = np.sqrt(3)*Lx # design domain longer side
22 Lex = Lx/Ns # element shorter side
23 Ley = np.sqrt(3)*Lex # element longer side
24 # Generate Mesh
25 coor, inci, etype, sym = get_mesh(Ns, Lex, Ley)
26 # check directories
27 rpath = '../dataset/SILP/'
28 if not os.path.exists(rpath):
29     print('missing SILP dataset')
30     sys.exit()
31 if not os.path.exists('./top_opt'):
32     os.mkdir('./top_opt')
33 if not os.path.exists('./top'):
34     os.mkdir('./top')
35 if not os.path.exists('./sen'):
36     os.mkdir('./sen')
37 if not os.path.exists('./dis'):
38     os.mkdir('./dis')
39 if not os.path.exists('./nu_Ey_vol'):
40     os.mkdir('./nu_Ey_vol')

```

The loop through the selected files is started. The input files indices, the input data and the pointers relating each input with the corresponding iterations of the optimization processes are read from the “fid.npy”, “inp.npy” and “ptr2opt.npy” files.

```

41 file = file_ini
42 while (file < file_lim) and (os.path.exists(rpath + 'f{:04d}'.format(file))):
43     ### Read files
44     print('> reading files of f{:04d}'.format(file))
45     # input files id, input data and pointers to optimization
46     if not os.path.exists(rpath + 'f{:04d}/fid.npy'.format(file)):
47         print('missing : f{:04d}/fid.npy'.format(file))
48         sys.exit()
49     list_fid = np.load(rpath + 'f{:04d}/fid.npy'.format(file))
50     if not os.path.exists(rpath + 'f{:04d}/inp.npy'.format(file)):
51         print('missing : f{:04d}/inp.npy'.format(file))
52         sys.exit()
53     list_inp = np.load(rpath + 'f{:04d}/inp.npy'.format(file))
54     if not os.path.exists(rpath + 'f{:04d}/ptr2opt.npy'.format(file)):
55         print('missing : f{:04d}/ptr2opt.npy'.format(file))
56         sys.exit()
57     list_ptr2opt = np.load(rpath + 'f{:04d}/ptr2opt.npy'.format(file))

```

According to the boolean flags, the optimized topologies, topologies, sensitivity vectors, displacements vectors, Poisson’s ratio values, Young’s modulus values and volume values are read from the “top_opt.npy”, “top.npy”, “dC00_0.npy”, “dC00_1.npy”, “dC00_2.npy”, “dC00_w.npy”, “dC11_0.npy”, “dC11_1.npy”, “dC11_2.npy”, “dC11_w.npy”, “dC22_0.npy”, “dC22_1.npy”, “dC22_2.npy”, “dC22_w.npy”, “dis_xx.npy”, “dis_yy.npy”, “dis_xy.npy”, “nu.npy”, “Ey.npy” and “vol.npy” files.

```

58 # optimized topology
59 if fig_top_opt:
60     if not os.path.exists(rpath + 'f{:04d}/top_opt.npy'.format(file)):
61         print('missing : f{:04d}/top_opt.npy'.format(file))
62         sys.exit()
63     list_top_opt = np.load(rpath + 'f{:04d}/top_opt.npy'.format(file))
64 # topology vectors
65 if fig_top or fig_dis:
66     if not os.path.exists(rpath + 'f{:04d}/top.npy'.format(file)):
67         print('missing : f{:04d}/top.npy'.format(file))
68         sys.exit()
69     list_top = np.load(rpath + 'f{:04d}/top.npy'.format(file))
70 # sensitivity vectors
71 if fig_sen:
72     if not os.path.exists(rpath + 'f{:04d}/dC00_0.npy'.format(file)):
73         print('missing : f{:04d}/dC00_0.npy'.format(file))
74         sys.exit()
75     list_dC00_0 = np.load(rpath + 'f{:04d}/dC00_0.npy'.format(file))
76     if not os.path.exists(rpath + 'f{:04d}/dC00_1.npy'.format(file)):
77         print('missing : f{:04d}/dC00_1.npy'.format(file))
78         sys.exit()
79     list_dC00_1 = np.load(rpath + 'f{:04d}/dC00_1.npy'.format(file))
80     if not os.path.exists(rpath + 'f{:04d}/dC00_2.npy'.format(file)):
81         print('missing : f{:04d}/dC00_2.npy'.format(file))
82         sys.exit()
83     list_dC00_2 = np.load(rpath + 'f{:04d}/dC00_2.npy'.format(file))

```

```

84     if not os.path.exists(rpath + 'f{:04d}/dC00_w.npy'.format(file)):
85         print('missing : f{:04d}/dC00_w.npy'.format(file))
86         sys.exit()
87     list_dC00_w = np.load(rpath + 'f{:04d}/dC00_w.npy'.format(file))
88     if not os.path.exists(rpath + 'f{:04d}/dC11_0.npy'.format(file)):
89         print('missing : f{:04d}/dC11_0.npy'.format(file))
90         sys.exit()
91     list_dC11_0 = np.load(rpath + 'f{:04d}/dC11_0.npy'.format(file))
92     if not os.path.exists(rpath + 'f{:04d}/dC11_1.npy'.format(file)):
93         print('missing : f{:04d}/dC11_1.npy'.format(file))
94         sys.exit()
95     list_dC11_1 = np.load(rpath + 'f{:04d}/dC11_1.npy'.format(file))
96     if not os.path.exists(rpath + 'f{:04d}/dC11_2.npy'.format(file)):
97         print('missing : f{:04d}/dC11_2.npy'.format(file))
98         sys.exit()
99     list_dC11_2 = np.load(rpath + 'f{:04d}/dC11_2.npy'.format(file))
100    if not os.path.exists(rpath + 'f{:04d}/dC11_w.npy'.format(file)):
101        print('missing : f{:04d}/dC11_w.npy'.format(file))
102        sys.exit()
103    list_dC11_w = np.load(rpath + 'f{:04d}/dC11_w.npy'.format(file))
104    if not os.path.exists(rpath + 'f{:04d}/dC22_0.npy'.format(file)):
105        print('missing : f{:04d}/dC22_0.npy'.format(file))
106        sys.exit()
107    list_dC22_0 = np.load(rpath + 'f{:04d}/dC22_0.npy'.format(file))
108    if not os.path.exists(rpath + 'f{:04d}/dC22_1.npy'.format(file)):
109        print('missing : f{:04d}/dC22_1.npy'.format(file))
110        sys.exit()
111    list_dC22_1 = np.load(rpath + 'f{:04d}/dC22_1.npy'.format(file))
112    if not os.path.exists(rpath + 'f{:04d}/dC22_2.npy'.format(file)):
113        print('missing : f{:04d}/dC22_2.npy'.format(file))
114        sys.exit()
115    list_dC22_2 = np.load(rpath + 'f{:04d}/dC22_2.npy'.format(file))
116    if not os.path.exists(rpath + 'f{:04d}/dC22_w.npy'.format(file)):
117        print('missing : f{:04d}/dC22_w.npy'.format(file))
118        sys.exit()
119    list_dC22_w = np.load(rpath + 'f{:04d}/dC22_w.npy'.format(file))
120    # displacements vectors
121    if fig_dis:
122        if not os.path.exists(rpath + 'f{:04d}/dis_xx.npy'.format(file)):
123            print('missing : f{:04d}/dis_xx.npy'.format(file))
124            sys.exit()
125        list_dis_xx = np.load(rpath + 'f{:04d}/dis_xx.npy'.format(file))
126        if not os.path.exists(rpath + 'f{:04d}/dis_yy.npy'.format(file)):
127            print('missing : f{:04d}/dis_yy.npy'.format(file))
128            sys.exit()
129        list_dis_yy = np.load(rpath + 'f{:04d}/dis_yy.npy'.format(file))
130        if not os.path.exists(rpath + 'f{:04d}/dis_xy.npy'.format(file)):
131            print('missing : f{:04d}/dis_xy.npy'.format(file))
132            sys.exit()
133        list_dis_xy = np.load(rpath + 'f{:04d}/dis_xy.npy'.format(file))
134    # Poisson's ratio, Young's modulus and volume
135    if fig_nu_Ey_vol:
136        if not os.path.exists(rpath + 'f{:04d}/nu.npy'.format(file)):
137            print('missing : f{:04d}/nu.npy'.format(file))
138            sys.exit()
139        list_nu = np.load(rpath + 'f{:04d}/nu.npy'.format(file))
140        if not os.path.exists(rpath + 'f{:04d}/Ey.npy'.format(file)):
141            print('missing : f{:04d}/Ey.npy'.format(file))
142            sys.exit()
143        list_Ey = np.load(rpath + 'f{:04d}/Ey.npy'.format(file))
144        if not os.path.exists(rpath + 'f{:04d}/vol.npy'.format(file)):
145            print('missing : f{:04d}/vol.npy'.format(file))
146            sys.exit()
147        list_vol = np.load(rpath + 'f{:04d}/vol.npy'.format(file))

```

If “fig_top_opt” is *True*, images of the optimized topologies are saved. The optimized topologies are represented as grayscale images, solid elements are represented in black and void elements are represented in gray.

```

148    """ Generate figures
149    print(' : generating figures')
150    # optimized topology
151    if fig_top_opt:
152        print(' : optimized topology...')
153        for k in range(len(list_fid)):
154            plt.figure(num=0).clear()
155            fig,ax = plt.subplots(num=0)
156            fid = list_fid[k]
157            x = list_top_opt[k]
158            x = np.unpackbits(x,axis=None).astype(float)
159            xt = np.ndarray((Nt),dtype=bool)
160            for k in range(N):
161                xt[sym[k,:]] = x[k]
162            polys = clct.PolyCollection(coor[inci],cmap='gray_r',edgecolor=(0,0,0,0))
163            polys.set_array(xt+1.0)
164            polys.set_clim(0.0,2.0)
165            ax.add_collection(polys)

```

```

166         ax.set_aspect('equal')
167         ax.set_xlim([-2*Lx,2*Lx])
168         ax.set_ylim([-Ly,Ly])
169         ax.axis('off')
170         fig.set_size_inches(8,7)
171         plt.savefig('..top_opt/f{:05d}.png'.format(fid),bbox_inches='tight',pad_inches=0.05,dpi=100)

```

If “fig_top” is *True*, images of the topology vectors are saved. The topologies are represented as grayscale images.

```

172     # topology vectors
173     if fig_top:
174         print(' : topology vectors...')
175         for k in range(len(list_fid)):
176             fid = list_fid[k]
177             j = 0
178             for kk in range(list_ptr2opt[k],list_ptr2opt[k+1]):
179                 plt.figure(num=0).clear()
180                 fig,ax = plt.subplots(num=0)
181                 x = list_top[kk]
182                 x = np.unpackbits(x,axis=None).astype(float)
183                 xt = np.ndarray((Nt),dtype=bool)
184                 for k in range(N):
185                     xt[sym[k,:]] = x[k]
186                 polys = clct.PolyCollection(coor[inci],cmap='gray_r',edgecolor=(0,0,0,0))
187                 polys.set_array(xt+1.0)
188                 polys.set_clim(0.0,2.0)
189                 ax.add_collection(polys)
190                 ax.set_aspect('equal')
191                 ax.set_xlim([-2*Lx,2*Lx])
192                 ax.set_ylim([-Ly,Ly])
193                 ax.axis('off')
194                 fig.set_size_inches(8,7)
195                 plt.savefig('..top/f{:05d}_{:03d}.png'.format(fid,j),bbox_inches='tight',pad_inches=0.05,dpi=100)
196                 j += 1

```

If “fig_sen” is *True*, images of the sensitivity vectors are saved. Each sensitivity map is represented as a single-channelled image, the “cividis” colormap is used instead of grayscale. An independent nonlinear scale is defined for each image, in order to improve contrast resolution, this scale is computed by the function “adjust.adjust”. The python script “adjust.py” is detailed in the next section.

The twelve sensitivity maps are saved in a single figure, in a 3×4 grid of subplots. The first row corresponds to sensitivity maps of C_{00} ; the second row to the sensitivity maps of C_{11} ; and the third row to the sensitivity maps of C_{22} . The first column corresponds to the CGS-0 approximations; the second column to the CGS-1 approximations; the third column to the CGS-2 approximations; and the fourth column to the exact sensitivity maps, obtained through WS approach.

```

197     # sensitivity vectors
198     if fig_sen:
199         print(' : sensitivity vectors...')
200         for k in range(len(list_fid)):
201             fid = list_fid[k]
202             j = 0
203             for kk in range(list_ptr2opt[k],list_ptr2opt[k+1]):
204                 plt.figure(num=0).clear()
205                 fig,ax = plt.subplots(nrows=3,ncols=4,num=0)
206                 xmin = -2*Lx
207                 xmax = 2*Lx
208                 ymin = -Ly
209                 ymax = Ly
210                 dC00_0 = list_dC00_0[kk]
211                 sens_plot = np.ndarray((Nt))
212                 adjusted = adjust(dC00_0,N)
213                 for k in range(N):
214                     sens_plot[sym[k,:]] = adjusted[k]
215                 polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
216                 polys.set_array(sens_plot)
217                 ax[0,0].add_collection(polys)
218                 ax[0,0].set_aspect('equal')
219                 ax[0,0].set_xlim([xmin,xmax])
220                 ax[0,0].set_ylim([ymin,ymax])
221                 ax[0,0].axis('off')
222                 dC00_1 = list_dC00_1[kk]
223                 sens_plot = np.ndarray((Nt))
224                 adjusted = adjust(dC00_1,N)
225                 for k in range(N):
226                     sens_plot[sym[k,:]] = adjusted[k]
227                 polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
228                 polys.set_array(sens_plot)
229                 ax[0,1].add_collection(polys)

```

```

230     ax[0,1].set_aspect('equal')
231     ax[0,1].set_xlim([xmin,xmax])
232     ax[0,1].set_ylim([ymin,ymax])
233     ax[0,1].axis('off')
234     dC00_2 = list_dC00_2[kk]
235     sens_plot = np.ndarray((Nt))
236     adjusted = adjust(dC00_2,N)
237     for k in range(N):
238         sens_plot[sym[k,:]] = adjusted[k]
239     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
240     polys.set_array(sens_plot)
241     ax[0,2].add_collection(polys)
242     ax[0,2].set_aspect('equal')
243     ax[0,2].set_xlim([xmin,xmax])
244     ax[0,2].set_ylim([ymin,ymax])
245     ax[0,2].axis('off')
246     dC00_w = list_dC00_w[kk]
247     sens_plot = np.ndarray((Nt))
248     adjusted = adjust(dC00_w,N)
249     for k in range(N):
250         sens_plot[sym[k,:]] = adjusted[k]
251     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
252     polys.set_array(sens_plot)
253     ax[0,3].add_collection(polys)
254     ax[0,3].set_aspect('equal')
255     ax[0,3].set_xlim([xmin,xmax])
256     ax[0,3].set_ylim([ymin,ymax])
257     ax[0,3].axis('off')
258     dC11_0 = list_dC11_0[kk]
259     sens_plot = np.ndarray((Nt))
260     adjusted = adjust(dC11_0,N)
261     for k in range(N):
262         sens_plot[sym[k,:]] = adjusted[k]
263     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
264     polys.set_array(sens_plot)
265     ax[1,0].add_collection(polys)
266     ax[1,0].set_aspect('equal')
267     ax[1,0].set_xlim([xmin,xmax])
268     ax[1,0].set_ylim([ymin,ymax])
269     ax[1,0].axis('off')
270     dC11_1 = list_dC11_1[kk]
271     sens_plot = np.ndarray((Nt))
272     adjusted = adjust(dC11_1,N)
273     for k in range(N):
274         sens_plot[sym[k,:]] = adjusted[k]
275     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
276     polys.set_array(sens_plot)
277     ax[1,1].add_collection(polys)
278     ax[1,1].set_aspect('equal')
279     ax[1,1].set_xlim([xmin,xmax])
280     ax[1,1].set_ylim([ymin,ymax])
281     ax[1,1].axis('off')
282     dC11_2 = list_dC11_2[kk]
283     sens_plot = np.ndarray((Nt))
284     adjusted = adjust(dC11_2,N)
285     for k in range(N):
286         sens_plot[sym[k,:]] = adjusted[k]
287     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
288     polys.set_array(sens_plot)
289     ax[1,2].add_collection(polys)
290     ax[1,2].set_aspect('equal')
291     ax[1,2].set_xlim([xmin,xmax])
292     ax[1,2].set_ylim([ymin,ymax])
293     ax[1,2].axis('off')
294     dC11_w = list_dC11_w[kk]
295     sens_plot = np.ndarray((Nt))
296     adjusted = adjust(dC11_w,N)
297     for k in range(N):
298         sens_plot[sym[k,:]] = adjusted[k]
299     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
300     polys.set_array(sens_plot)
301     ax[1,3].add_collection(polys)
302     ax[1,3].set_aspect('equal')
303     ax[1,3].set_xlim([xmin,xmax])
304     ax[1,3].set_ylim([ymin,ymax])
305     ax[1,3].axis('off')
306     dC22_0 = list_dC22_0[kk]
307     sens_plot = np.ndarray((Nt))
308     adjusted = adjust(dC22_0,N)
309     for k in range(N):
310         sens_plot[sym[k,:]] = adjusted[k]
311     polys = clct.PolyCollection(coor[inci],cmap='cividis',edgecolor=(0,0,0,0))
312     polys.set_array(sens_plot)
313     ax[2,0].add_collection(polys)
314     ax[2,0].set_aspect('equal')
315     ax[2,0].set_xlim([xmin,xmax])
316     ax[2,0].set_ylim([ymin,ymax])
317     ax[2,0].axis('off')
318     dC22_1 = list_dC22_1[kk]
319     sens_plot = np.ndarray((Nt))
320     adjusted = adjust(dC22_1,N)

```

```

321         for k in range(N):
322             sens_plot[sym[k,:]] = adjusted[k]
323             polys = clct.PolyCollection(coor[inci], cmap='cividis', edgecolor=(0,0,0,0))
324             polys.set_array(sens_plot)
325             ax[2,1].add_collection(polys)
326             ax[2,1].set_aspect('equal')
327             ax[2,1].set_xlim([xmin,xmax])
328             ax[2,1].set_ylim([ymin,ymax])
329             ax[2,1].axis('off')
330             dC22_2 = list_dC22_2[kk]
331             sens_plot = np.ndarray((Nt))
332             adjusted = adjust(dC22_2,N)
333             for k in range(N):
334                 sens_plot[sym[k,:]] = adjusted[k]
335                 polys = clct.PolyCollection(coor[inci], cmap='cividis', edgecolor=(0,0,0,0))
336                 polys.set_array(sens_plot)
337                 ax[2,2].add_collection(polys)
338                 ax[2,2].set_aspect('equal')
339                 ax[2,2].set_xlim([xmin,xmax])
340                 ax[2,2].set_ylim([ymin,ymax])
341                 ax[2,2].axis('off')
342                 dC22_w = list_dC22_w[kk]
343                 sens_plot = np.ndarray((Nt))
344                 adjusted = adjust(dC22_w,N)
345                 for k in range(N):
346                     sens_plot[sym[k,:]] = adjusted[k]
347                     polys = clct.PolyCollection(coor[inci], cmap='cividis', edgecolor=(0,0,0,0))
348                     polys.set_array(sens_plot)
349                     ax[2,3].add_collection(polys)
350                     ax[2,3].set_aspect('equal')
351                     ax[2,3].set_xlim([xmin,xmax])
352                     ax[2,3].set_ylim([ymin,ymax])
353                     ax[2,3].axis('off')
354                     fig.set_size_inches(20,13)
355                     plt.savefig('./sen/f{:05d}_{:03d}.png'.format(fid,j), bbox_inches='tight', pad_inches=0.05, dpi=100)
356                     j += 1

```

If “fig_dis” is *True*, images of the deformed cell are saved, considering the three imposed macro-displacements. In order to visualize the displacements field, the topologies are represented in the deformed mesh. The scale is kept the same for all figures, so the influence of each topological change can be perceived.

The three displacements fields are saved in a single figure, in a 1×3 grid of subplots. The first column corresponds to the deformed cell when \hat{u}_{xx} is imposed; the second column to the deformed cell when \hat{u}_{yy} is imposed; and the third column to the deformed cell when \hat{u}_{xy} is imposed.

```

357     # displacements vectors
358     if fig_dis:
359         print(' : displacements vectors...')
360         for k in range(len(list_fid)):
361             fid = list_fid[k]
362             j = 0
363             scale = 0.10
364             xmin = -2*Lx*(1.05+scale)
365             xmax = 2*Lx*(1.05+scale)
366             ymin = -Ly*(1.15+scale)
367             ymax = Ly*(1.15+scale)
368             for kk in range(list_ptr2opt[k], list_ptr2opt[k+1]):
369                 plt.figure(num=0).clear()
370                 fig, ax = plt.subplots(nrows=1, ncols=3, num=0)
371                 x = list_top[kk]
372                 x = np.unpackbits(x, axis=None).astype(float)
373                 xt = np.ndarray((Nt), dtype=bool)
374                 for k in range(N):
375                     xt[sym[k,:]] = x[k]
376                     dis_xx = list_dis_xx[kk]
377                     umat = np.reshape(dis_xx, coor.shape)
378                     coor_dis = coor + scale*umat
379                     polys = clct.PolyCollection(coor_dis[inci], cmap='gray_r', edgecolor=(0,0,0,0))
380                     polys.set_array(xt+1.0)
381                     polys.set_clim(0.0, 2.0)
382                     ax[0].add_collection(polys)
383                     ax[0].set_aspect('equal')
384                     ax[0].set_xlim([xmin,xmax])
385                     ax[0].set_ylim([ymin,ymax])
386                     ax[0].axis('off')
387                     dis_yy = list_dis_yy[kk]
388                     umat = np.reshape(dis_yy, coor.shape)
389                     coor_dis = coor + scale*umat
390                     polys = clct.PolyCollection(coor_dis[inci], cmap='gray_r', edgecolor=(0,0,0,0))
391                     polys.set_array(xt+1.0)
392                     polys.set_clim(0.0, 2.0)
393                     ax[1].add_collection(polys)
394                     ax[1].set_aspect('equal')
395                     ax[1].set_xlim([xmin,xmax])
396                     ax[1].set_ylim([ymin,ymax])
397                     ax[1].axis('off')

```



```

398         dis_xy = list_dis_xy[kk]
399         umat = np.reshape(dis_xy,coor.shape)
400         coor_dis = coor + scale*umat
401         polys = clct.PolyCollection(coor_dis[inci],cmap='gray_r',edgecolor=(0,0,0,0))
402         polys.set_array(xt+1.0)
403         polys.set_clim(0.0,2.0)
404         ax[2].add_collection(polys)
405         ax[2].set_aspect('equal')
406         ax[2].set_xlim([xmin,xmax])
407         ax[2].set_ylim([ymin,ymax])
408         ax[2].axis('off')
409         fig.set_size_inches(20,8)
410         fig.savefig('./dis/f{:05d}_{:03d}.png'.format(fid,j),bbox_inches='tight',pad_inches=0,dpi=100)
411         j += 1

```

If “fig_nu_Ey_vol” is *True*, plots of Poisson’s ratio, Young’s modulus and volume are saved. The evolution of these functions throughout each optimization procedure is plotted in the same figure, in a 3×1 grid of subplots.

The file counter “file” is updated so the next images can be generated.

```

412 # Poisson's ratio, Young's modulus and volume
413 if fig_nu_Ey_vol:
414     print(' : Poisson\'s ratio, Young\'s modulus and volume...')
415     for k in range(len(list_fid)):
416         plt.figure(num=0).clear()
417         fig,ax = plt.subplots(nrows=3,ncols=1,num=0)
418         fid = list_fid[k]
419         inp = list_inp[k]
420         Eymin = inp[1]
421         nu = list_nu[list_ptr2opt[k]:list_ptr2opt[k+1]]
422         Ey = list_Ey[list_ptr2opt[k]:list_ptr2opt[k+1]]
423         vol = list_vol[list_ptr2opt[k]:list_ptr2opt[k+1]]
424         size = len(nu)
425         delta = max(nu)-min(nu)
426         miny = min(nu)-0.02*delta
427         maxy = max(nu)+0.02*delta
428         ax[0].plot(nu,'ok-',linewidth=2)
429         ax[0].axis([-0.75, size-0.25, miny, maxy])
430         ax[0].set_ylabel('Poisson\'s ratio',fontsize=18)
431         ax[0].grid()
432         delta = max(Ey)-min(Ey)
433         miny = Eymin-0.02*delta
434         maxy = max(Ey)+0.02*delta
435         ax[1].plot(Ey,'ok-',linewidth=2)
436         ax[1].plot([-0.75, size-0.25], [Eymin,Eymin], 'k--',linewidth=2)
437         ax[1].axis([-0.75, size-0.25, miny, maxy])
438         ax[1].set_ylabel('Young\'s modulus [Pa]',fontsize=18)
439         ax[1].grid()
440         delta = max(vol)-min(vol)
441         miny = min(vol)-0.02*delta
442         maxy = max(vol)+0.02*delta
443         ax[2].plot(vol,'ok-',linewidth=2)
444         ax[2].axis([-0.75, size-0.25, miny, maxy])
445         ax[2].set_ylabel('volume fraction',fontsize=18)
446         ax[2].grid()
447         ax[2].set_xlabel('iteration',fontsize=18)
448         fig.set_size_inches(8,13)
449         fig.savefig('./nu_Ey_vol/f{:05d}.png'.format(fid),bbox_inches='tight',pad_inches=0.05,dpi=100)
450     # prepare to read next file
451     file = file + 1

```

When all selected images are generated, the figure window is closed and the program terminates.

```

452 plt.close(fig=0)
453 print('done!')

```

3.5.2 ./sample/SILP/adjust.py

This script is used to adjust the sensitivity maps in order to improve contrast resolution of the generated figures. Firstly, the numpy module is imported.

```

1 import numpy as np

```

The “adjust” function is defined. The variable “delta” stores the difference between the maximal and minimal sensitivity values. The vector is shifted so that its median value becomes 0.0. The positive values are shifted up, and the nonpositive values are shifted down (the shift value is given by a fraction of “delta”, defined

by the variable “coef”). The absolute values of the negative part are taken, then, the logarithms of both parts are computed. Lastly, the former negative part is reflected over the horizontal axis that crosses its minimal value (“mval”). After performing all these procedures, the curve described by the sorted adjusted vector is considered. The variable “coef” is calibrated in so that the slope in the middle of the curve (“slope”) be similar to its mean slope (“ref”).

```

2 def adjust(sens,N):
3     slope = 1
4     ref = 0
5     coef = 1e-12
6     while slope > ref:
7         adjusted = sens.copy()
8         delta = max(adjusted)-min(adjusted)
9         adjusted = adjusted - np.median(adjusted)
10        mask = (adjusted > 0.0)
11        adjusted[mask] = adjusted[mask] + coef*delta
12        adjusted[~mask] = adjusted[~mask] - coef*delta
13        adjusted[mask] = np.log(adjusted[mask])
14        adjusted[~mask] = np.log(-adjusted[~mask])
15        mval = min(adjusted[~mask])
16        adjusted[~mask] = 2*mval - adjusted[~mask]
17        adj_sorted = np.sort(adjusted)
18        slope = (adj_sorted[N//2+2]-adj_sorted[N//2-3])/5
19        ref = (max(adjusted[mask])-min(adjusted[~mask]))/(N-1)
20        coef = 2*coef
21    return adjusted

```

3.6 Validation Procedure

The validation procedure consists mainly in independent, alternative implementations to perform each task of the optimization program that generates the dataset.

The validation codes are provided in the folder “./validation”. The bash script “metaval.sh” can be executed in order to build the required Cython codes. The script “metamaterial_val.py” can be executed to perform the following validations.

Firstly, four different topology vectors are created. The first one corresponds to a fully solid structure; the second one corresponds to a fully void structure; the third one corresponds to the initial topology used in the optimization procedures; and the fourth one corresponds to a random structure.

The homogenization procedure is performed for all the topologies. Then, the program checks if: the obtained properties are isotropic; the loads over opposing edges are anti-symmetric; the mean stress corresponds to the obtained elasticity matrix applied on the mean strain of the base cell; the mechanical properties are within their minimal and maximal possible values. The resultant loads are shown in a figure, so that it can be seen that there is no external load in the interior of the base cell (all nonzero loads are placed over the edges), and that the loads are anti-symmetric. The range of the loads (minimal and maximal values) are presented above each figure.

The “topopt.update” function is tested: the first topology is updated to be equal the fourth one; then, the resulting displacements and stiffness matrices are compared with the ones previously computed for the fourth topology. The error due to the shift maneuver (used to preserve the nonzero pattern of the stiffness matrix) is evaluated.

The fourth topology is used to validate the sensitivity values. The CGS values computed in the “silp_sens.pyx” script are compared with values obtained using estimations for the displacements vectors after altering each element, using the Conjugate Gradient Method. The WS values computed in the “topopt.py” script are compared with values obtained through a naive approach, using exact values for the displacements vectors after altering each element. The signs of the sensitivity values of the diagonal terms of \mathbf{C} are verified, as well as their monotonic behavior, as more steps are considered in the CGS approach. The error of the sensitivity values for the Poisson’s ratio, Young’s modulus and objective function are evaluated. These errors are plotted for CGS-0, CGS-1 and CGS-2, so it can be seen that, for most cases, CGS-2 is substantially more accurate than CGS-0.

Lastly, through visual verification, qualitative validations are performed for the conical filter used to smooth the sensitivity maps; for the morphological operators; and for the island removal procedure. To validate the smoothing filter, the fourth topology and a toy example are filtered, so it can be verified if the smoothing procedure is working as expected. To validate the morphological operator, the erosion, dilation, opening (erosion then dilation) and closing (dilation then erosion) operators are applied to the fourth topology. To validate the island removal procedure, it is performed for the third topology, the fourth topology and a toy example, the

connected and disconnected parts are shown in separate figures.

After executing the script “metamaterial_val.py”, a number of samples was selected from the generated dataset and visually verified, using the presented “sample.py” script. Some of them are shown in [section 4](#).

Furthermore, all input-output logs were checked, in order to verify if reasonable values were obtained for the mechanical properties of each optimized topology. It was verified if every pair of redundant data truly corresponds to the same value, that is, if each array “dC00_w.npy” is equal to the corresponding array “dC11_w.npy”. All observed results are coherent and indicate that the optimization program was properly implemented.

3.7 Unfixed Bug

The removal of disconnected solids from the optimized structure may yield an unsuitable solution. This bug is reported in the “Issues” tab of the github repository.

After concluding the current optimization process, all remaining disconnected solid elements are removed from the optimized topology. In the current version of the program, the final optimized topology is stored without verifying if the removal of disconnected solids resulted in a worse value for the objective function, or if it resulted in a structure that does not respect the constraint function. In cases that different parts of the structure are only connected by single nodes (not by edges), this approach may result in unsuitable structures.

To solve this issue (in a future version), all topologies can be stored, ordered from the best one obtained thus far to the worst one. Then, when an undesirable result is obtained, the next candidate can be taken and evaluated.

This issue has affected only a small number of optimized solutions. Less than 1% of the optimized solutions, stored in the ‘top_opt.npy’ files, break the constraint over the Young’s modulus. In the current version, suitable optimized solutions can be recovered using the data stored in the ‘top.npy’, ‘nu.npy’ and ‘Ey.npy’ files. As expected, the constraint is respected for 100% of the topologies from all iterations before the final removal of disconnected solids, stored in the ‘top.npy’ files.

4 Samples

The dataset stores everything generated in the optimization procedures for each one of the 18 382 unique pairs (ν^*, E_{\min}) .

To illustrate the size of the dataset, the homogenized mechanical properties from all the ‘nu.npy’ and ‘Ey.npy’ files, corresponding to topologies stored in all the ‘top.npy’ files, are shown in [Figure 7](#). There are 1 374 656 topologies stored in these files, however, around half of them have very similar (or identical) homogenized mechanical properties to other topologies from the dataset. After removing the duplicates, 616 862 topologies with unique mechanical properties are obtained, these are the ones shown in the scatter plot.

It can be seen that topologies with Young’s moduli near 100% have Poisson’s ratios near 0.30, which is the property of the base material. The range of the Poisson’s ratios increases as lower Young’s moduli are considered. When the Young’s moduli are near 0%, the obtained Poisson’s ratios range from nearly -1.0 to nearly 1.0 . There is a region with sparser points that corresponds to Poisson’s values between 0.20 and 0.40. It occurs because all optimizations start from a homogenized Poisson’s ratio of around 0.30 and it is desired to obtain values that are either higher than 0.40, or lower than 0.20. So, topologies move into this region only when undesirable steps are made in the optimization procedures. There are two regions with denser points around the minimal and the maximal Poisson’s values for each Young’s modulus. They occur because, for each considered minimal Young’s modulus (E_{\min}) the target Poisson’s values (ν^*) range from -1.0 to 1.0 . So, when E_{\min} is too high to allow extreme Poisson’s ratios, all optimizations that were not able to achieve their target values yield results near the minimal or maximal possible Poisson’s ratios for that Young’s modulus constraint.

The points corresponding to the optimized results, stored in all the ‘top_opt.npy’, ‘nu_opt.npy’ and ‘Ey_opt.npy’ files, are presented in [Figure 8](#). Once again, topologies with very similar (or identical) homogenized mechanical properties are disregarded: only around half of the 18 382 optimized topologies are shown in the scatter plot. The same observations can be made: there is an empty region corresponding to the Poisson’s values between 0.20 and 0.40 (unsearched region); and denser regions around the minimal and maximal Poisson’s values (just above the lower unreached region and just below the upper unreached region).

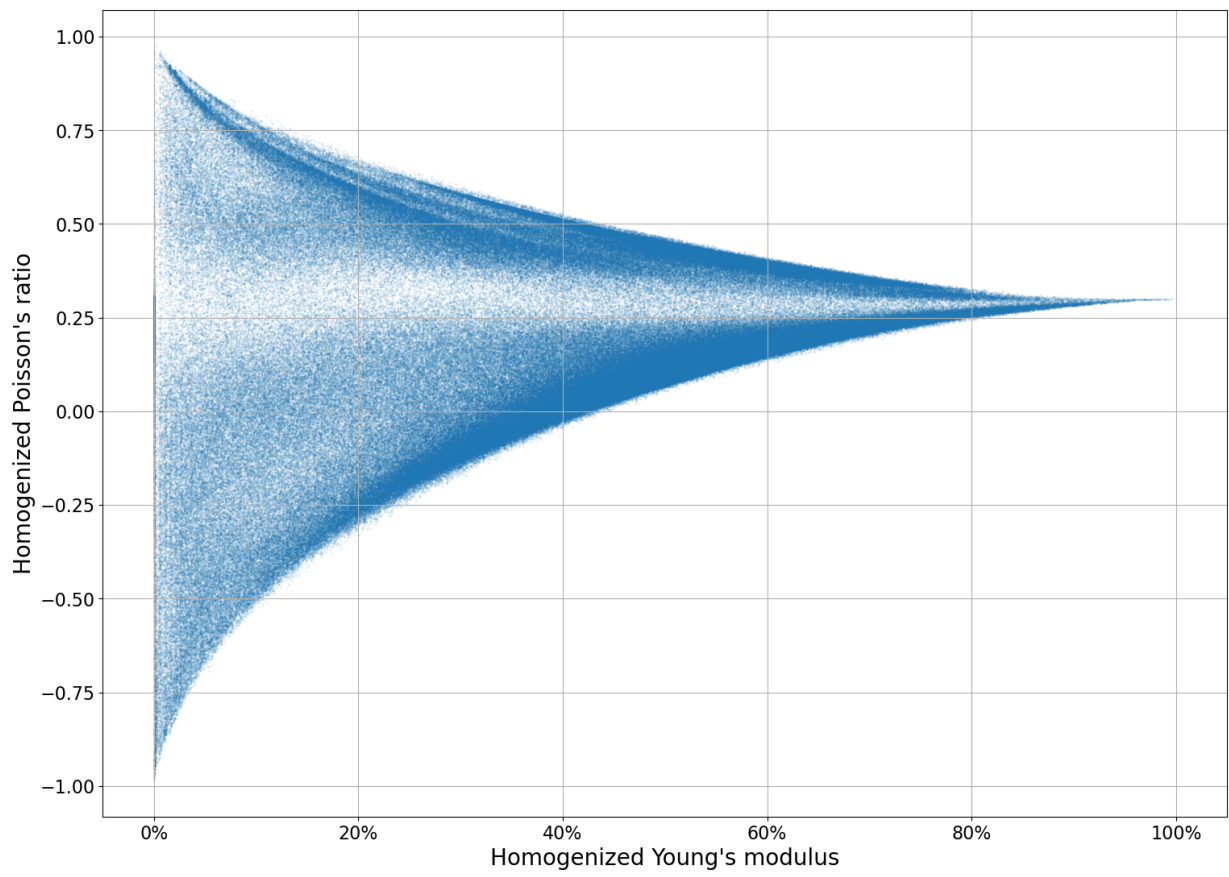


Figure 7: Properties of all generated metamaterials

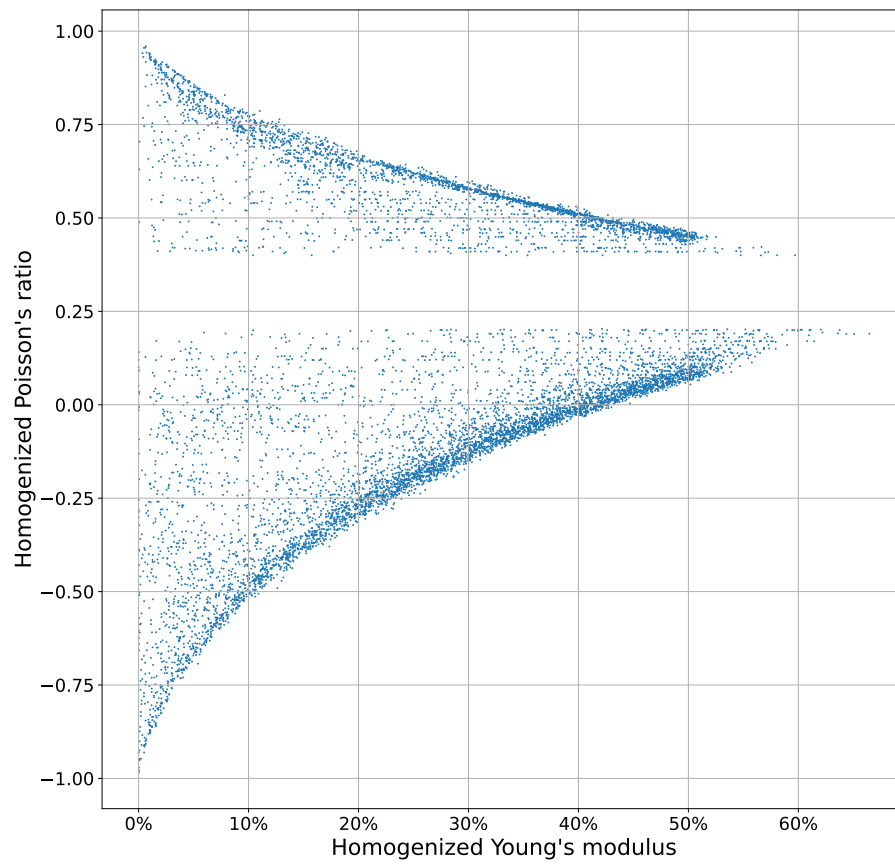


Figure 8: Properties of the optimized metamaterials

In **Figure 9**, a small sample of 19 optimized topologies is shown, the positions of the presented structures indicate their corresponding homogenized Poisson's ratios and Young's moduli. The solid part of the structures is represented in dark gray and the void part is represented in light gray.

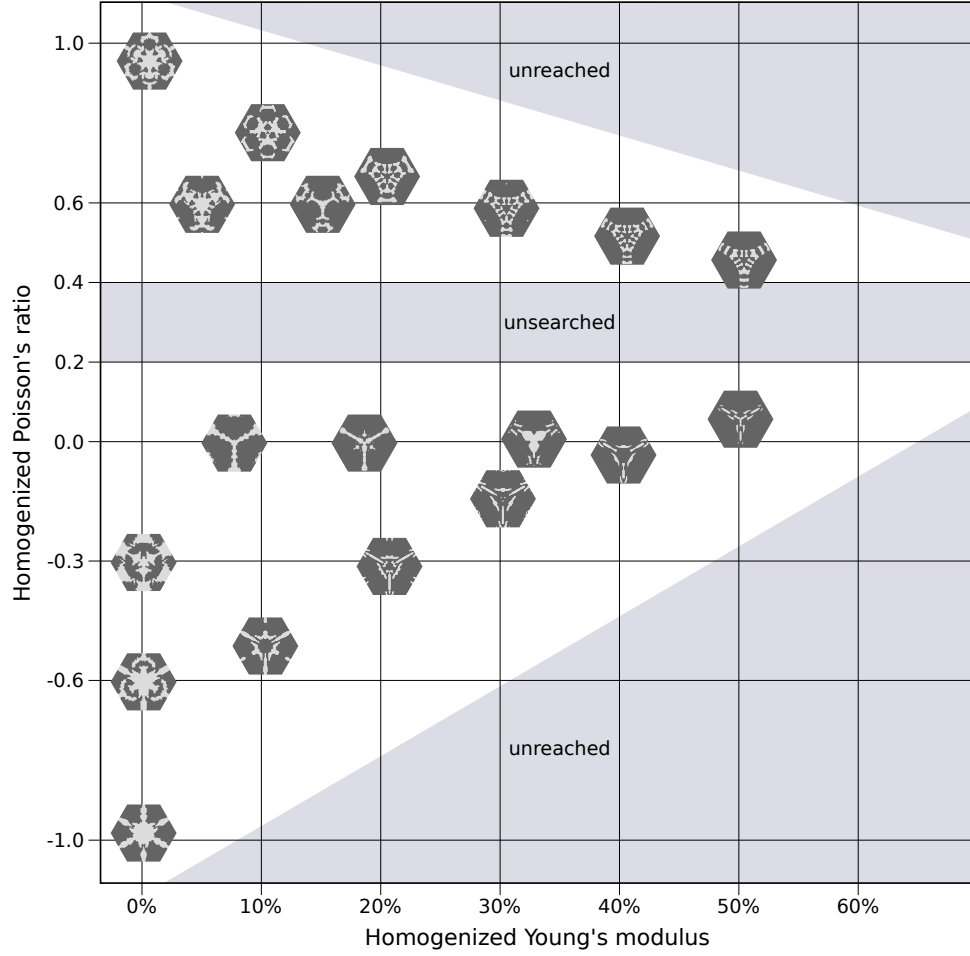


Figure 9: Optimized base cells and their homogenized properties

More samples are taken from the slices highlighted in **Figure 10**.

In the slice 'A', topologies with Young's moduli between 10% and 15% are taken, a sample of 28 structures with different Poisson's ratios is shown in **Figure 11**. Steps of 0.04 are considered. Topologies with Poisson's ratios ranging from -0.49 to 0.19 and from 0.41 to 0.77 are shown. The first row begins in lower unreached region and the second row ends in the upper unreached region. In the second row, the unsearched region corresponds to Poisson's ratios ranging from 0.20 to 0.40 .

In the slice 'B', topologies with Poisson's ratios between -0.02 and 0.02 are taken, a sample of 10 structures with different Young's moduli is shown in **Figure 12**. Steps of at most 7.0% are considered. Topologies with Young's moduli ranging from 0.0% to 44.0% are shown. In **Figure 10**, it can be seen that for the considered values of Poisson's ratios, Young's moduli above 45% correspond to the lower unreached region.

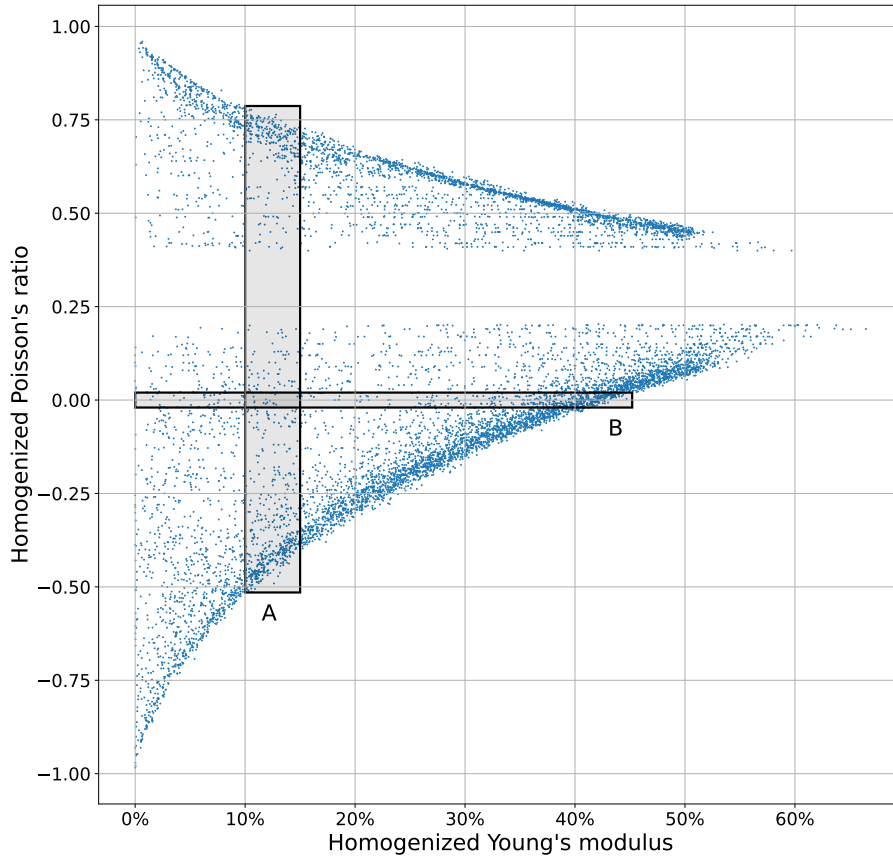


Figure 10: Slices of the scatter plot of optimized results

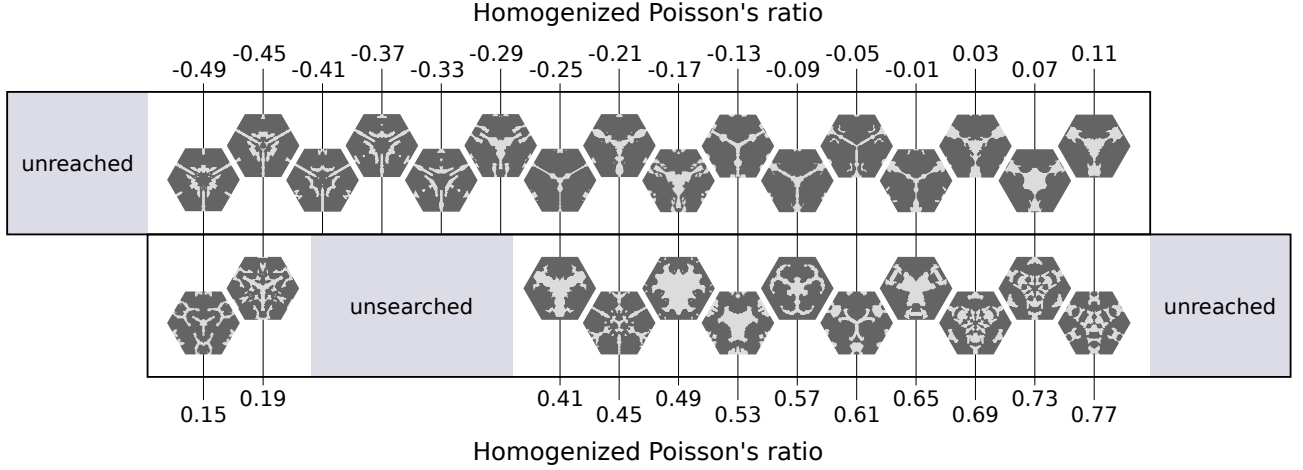


Figure 11: Optimized topologies with different Poisson's ratios (slice 'A')

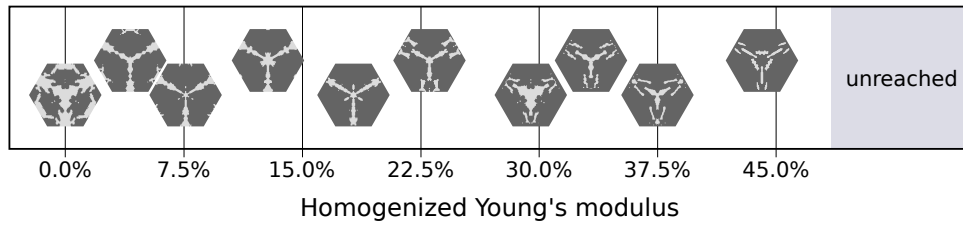


Figure 12: Optimized topologies with different Young's moduli (slice 'B')

To illustrate all the content stored in the dataset, the case with $\nu^* = -0.30$ and $E_{\min} = 20.0\%$ is considered. The obtained results are presented in Figures 13, 14, 15, 16 and 17, plotted by the “sample” script.

From Figure 13, it can be noted that the Poisson's ratio steadily improves when the Young's modulus is far from its minimal value. When the constraint is activated, the process become less stable because each time the solution of the linearized subproblem breaks the Young's modulus constraint, the structure is dilated. The dilation operations can be identified as abrupt increases in the volume fraction. The best obtained result corresponds to iteration 48, shown in Figure 14, its homogenized properties are $\hat{\nu} = -0.29$ and $\hat{E} = 20.4\%$. In Figure 15, all evaluated topologies are presented. Although a patience parameter of 30 was used, it took 81 iterations to conclude the procedure because the topology from iteration 51 has a smaller volume fraction than the 48-th topology, which resulted in a lower value for the objective function with volume penalization.

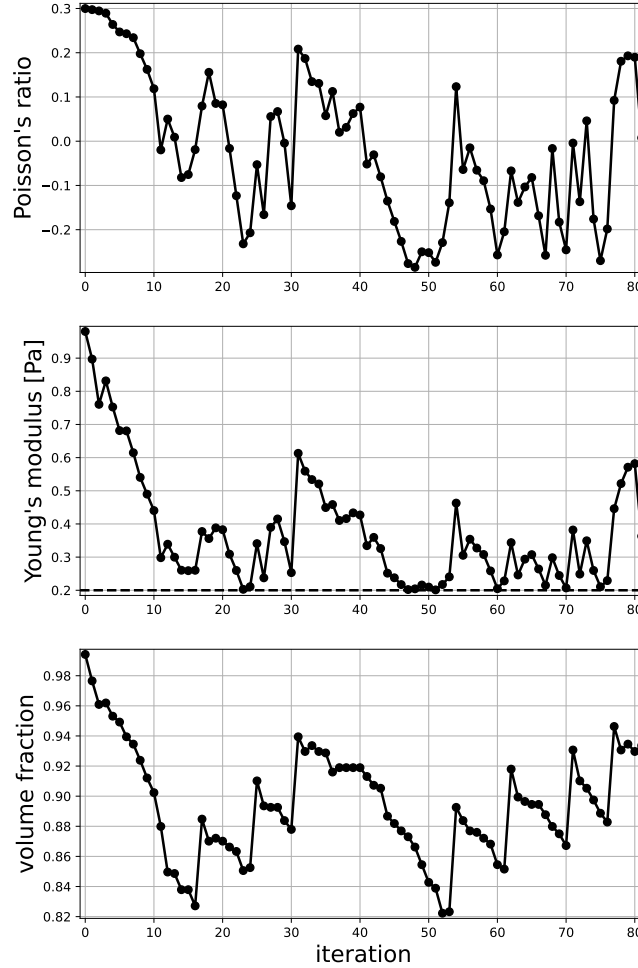


Figure 13: Poisson's ratio, Young's modulus and volume fraction for $\nu^* = -0.30$ and $E_{\min} = 20.0\%$



Figure 14: Optimized topology for $\nu^* = -0.30$ and $E_{\min} = 20.0\%$ (iteration 48)

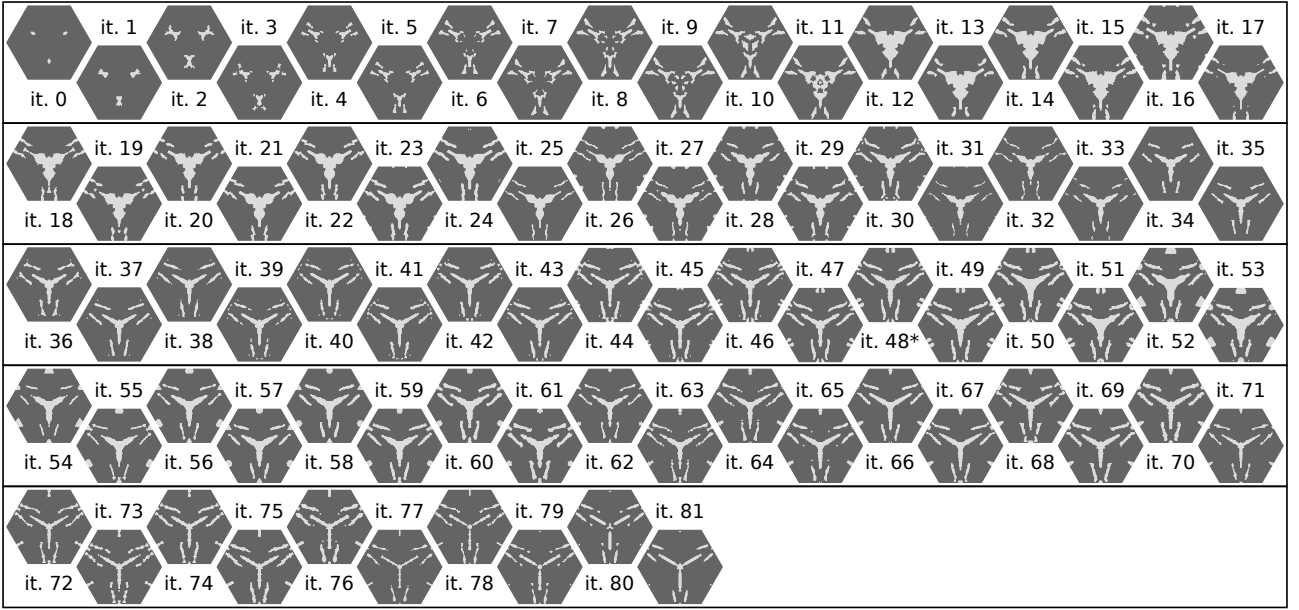


Figure 15: Topologies generated for $\nu^* = -0.30$ and $E_{\min} = 20.0\%$

Besides the topologies, homogenized properties and volume values, sensitivity and displacements vectors of each iteration are stored. **Figure 16** shows the 12 sensitivity vectors stored for the optimized topology. The maps for ΔC_{00} are in the first row, the ones for ΔC_{11} are in the second row and the ones for ΔC_{22} are in the third row. The first column corresponds to the CGS-0 estimations, the second column to the CGS-1 estimations, the third column to the CGS-2 estimations and the fourth column to the exact values, computed through WS approach. An independent nonlinear scale is adjusted for each one of the 12 maps, in order to improve contrast resolution, so what is being shown is only how the sensitivity values are distributed among the different elements, not their quantitative values. Although they are referred to as sensitivity values, they correspond to the **variations** of the diagonal terms of \mathbf{C} when the state of each augmented element is switched. To obtain proper sensitivity values, the signs of the values corresponding to solid augmented elements would have to be reversed.

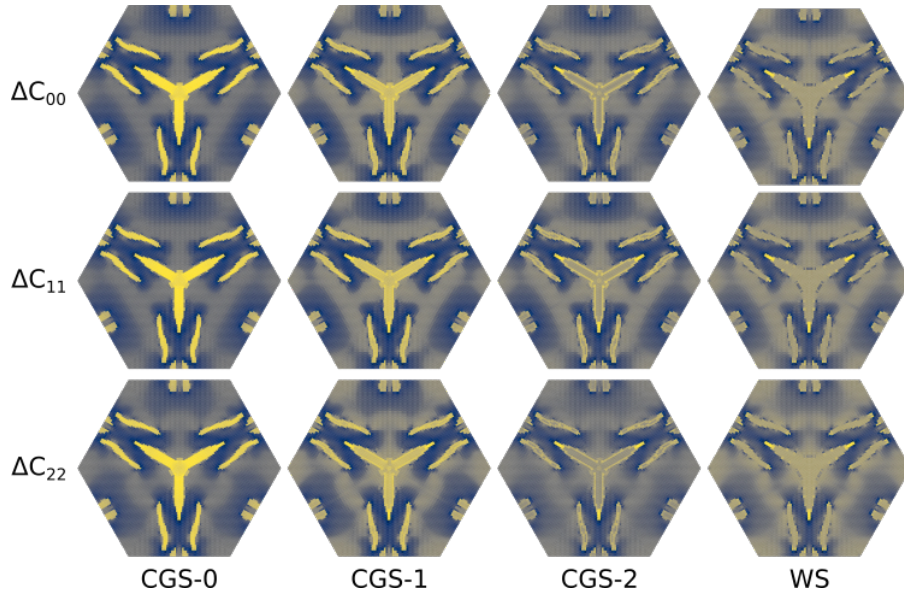


Figure 16: Sensitivity maps for the optimized topology (iteration 48)

In **Figure 17**, the deformed cell is presented for the three different values of macro-displacements imposed to perform the homogenization procedure. The displacements values are rescaled to improve visualization.

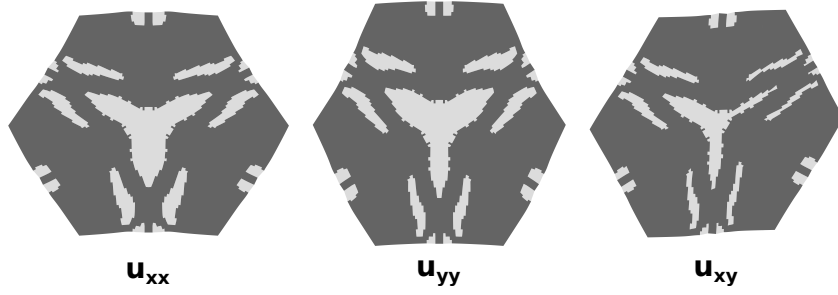


Figure 17: Deformed structures for the optimized topology (iteration 48)

To understand why there are cases in which a large number of iterations is necessary to achieve the stopping criterion, the case with maximal number of iterations is presented. The case in which $\nu^* = 0.57$ and $E_{\min} = 6.5\%$ needed 292 iterations to converge. **Figure 18** presents the evolution of the Poisson's ratio, Young's modulus and volume fraction over the optimization and **Figure 19** presents the topologies for some iterations. The best obtained result corresponds to iteration 262, its homogenized properties are $\hat{\nu} = 0.57$ and $\hat{E} = 15.3\%$.

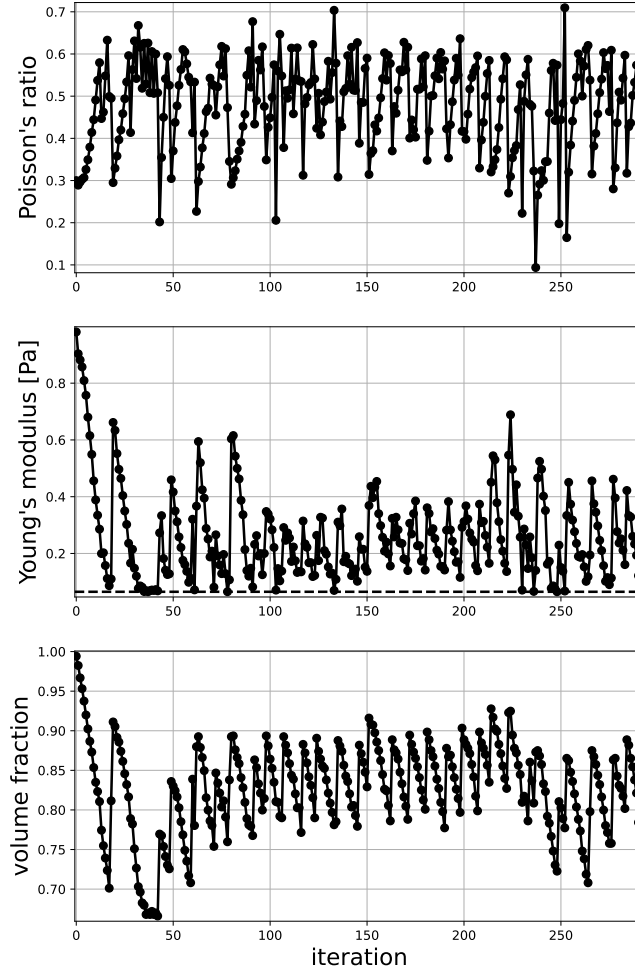


Figure 18: Poisson's ratio, Young's modulus and volume fraction for $\nu^* = 0.57$ and $E_{\min} = 6.5\%$

The procedure took so long because the dilation procedure often produced topologies that were in the basins of attraction of different local minima. Thus, more candidates were tried out throughout the iterative procedure. After iteration 262, the method was not able to keep improving the objective function, so the stopping criterion was reached.

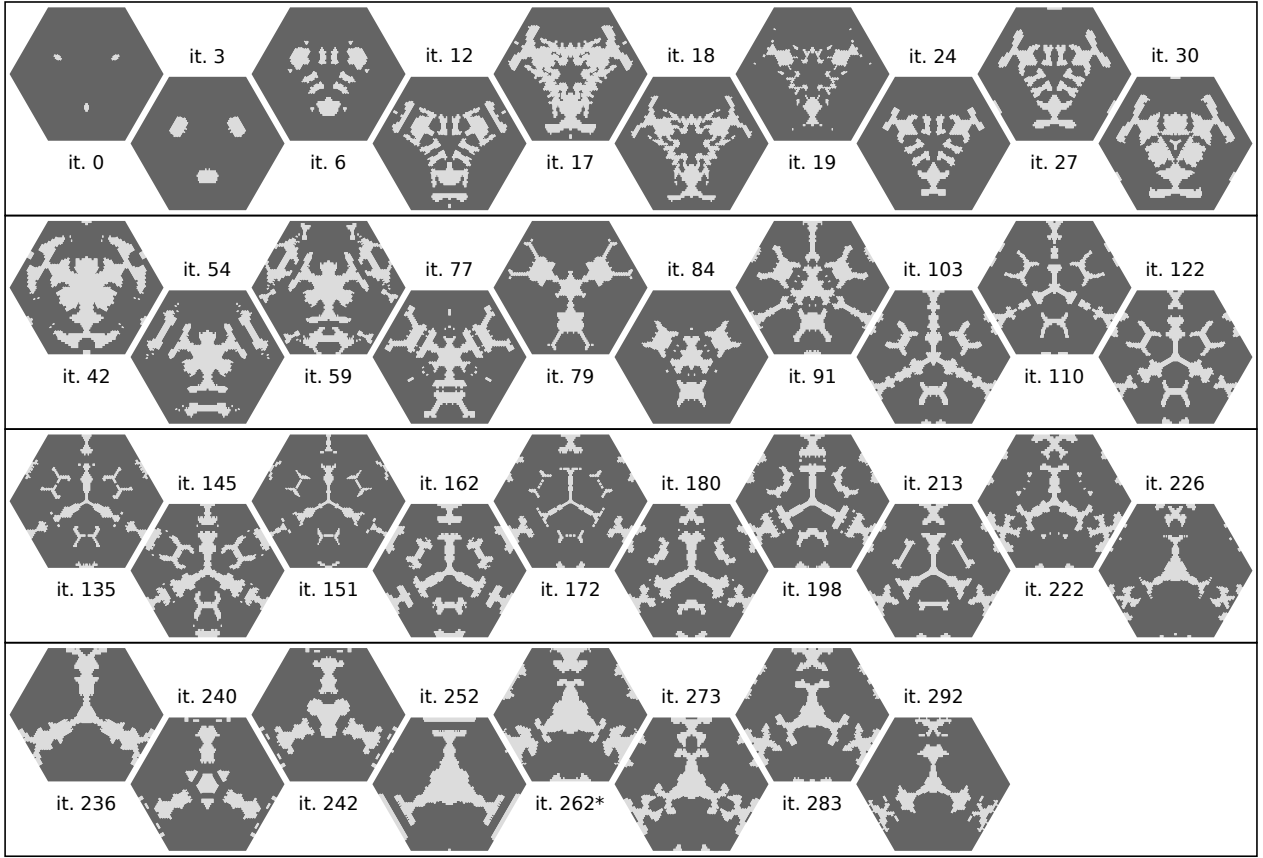


Figure 19: Topologies generated for $\nu^* = 0.57$ and $E_{\min} = 6.5\%$

Although it may be computationally expensive to perform so many iterations, in the SILP approach with the considered patience stopping criterion, a large number of iterations means that the method consistently improved the result along all the optimization procedure. Whenever there are sufficient time and computational resources, a more extensive exploration of the domain of possible solutions is advantageous, since it can yield more effective optimized structures.

On the other hand, there are cases in which a small number of iterations is enough to achieve the stopping criterion, the case with minimal number of iterations is presented. The case in which $\nu^* = -0.89$ and $E_{\min} = 40.5\%$ needed 37 iterations to converge. [Figure 20](#) presents the evolution of the Poisson's ratio, Young's modulus and volume fraction over the optimization and [Figure 21](#) presents the topologies for all iterations. The best obtained result corresponds to iteration 7, its homogenized properties are $\hat{\nu} = 0.08$ and $\hat{E} = 42.9\%$.

The high value for the minimal Young's modulus makes it easier to break the constraint, so only 7 iterations are performed before the first dilation. In this case, the perturbations from the dilation procedures were not enough to take the topologies out of the basins of attraction of the current local minimum. So, before each dilation, similar structures were obtained. Here, the first candidate ended up being better than all others obtained in subsequent iterations, so the optimization process took only 37 iterations to be concluded.

This behavior is undesirable, since the program is prevented from properly exploring the domain of possible solutions. In future versions, the stopping criterion may be improved to avoid such early convergences. Nonetheless, the obtained result is reasonable and suitable for the proposed dataset.

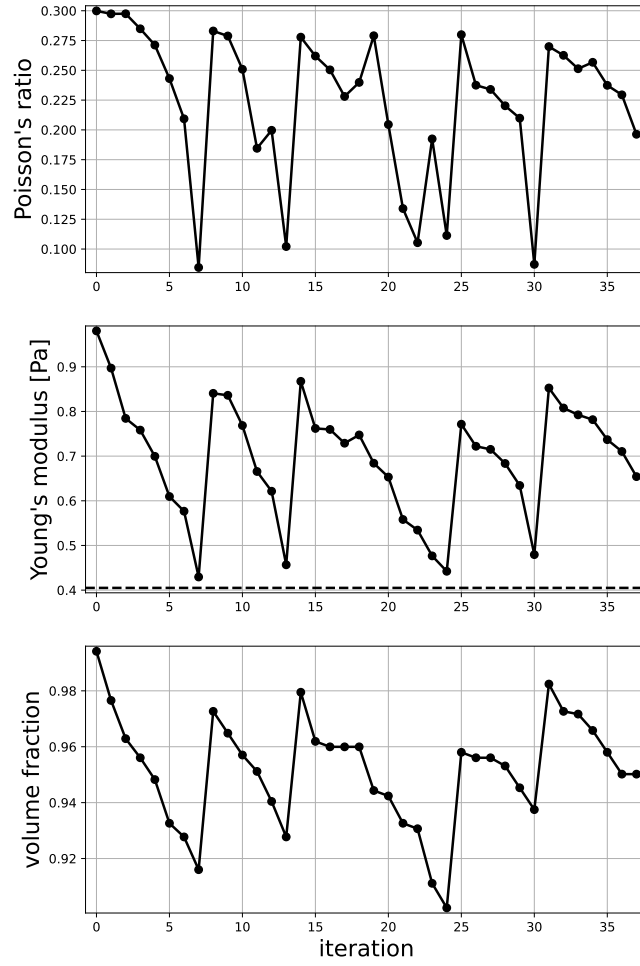


Figure 20: Poisson's ratio, Young's modulus and volume fraction for $\nu^* = -0.89$ and $E_{\min} = 40.5\%$

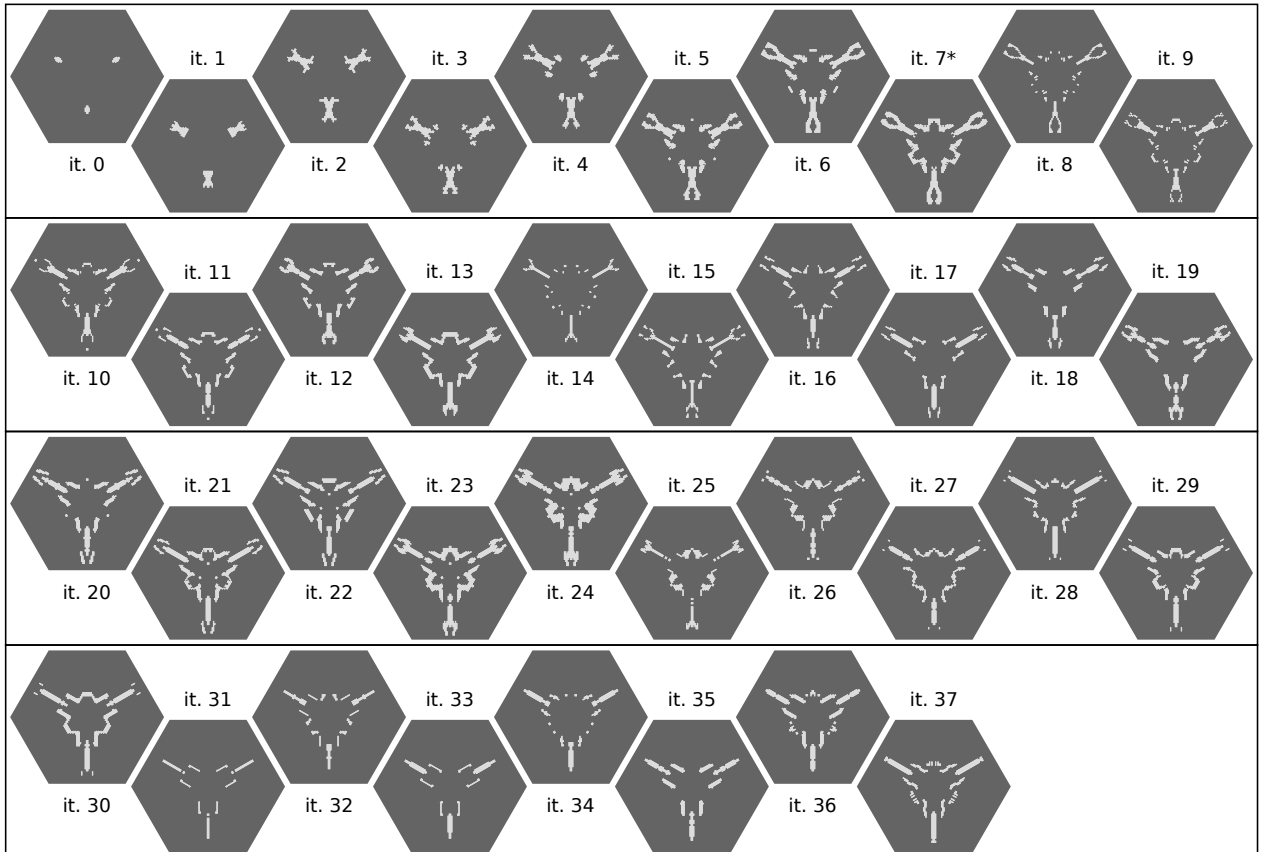


Figure 21: Topologies generated for $\nu^* = -0.89$ and $E_{\min} = 40.5\%$

It should be noted that there was a bug (already fixed) in the programs used to generate the data shown in this section. It did not compromise anything but the stability of the optimization procedures, so the presented samples are legitimate data. This only means that different results (possibly better) may be obtained when generating the dataset with the current version of the programs (presented in this document).

5 Summary

The specified values for each fixed parameter of the considered problem are presented in [Table 1](#).

Table 1: Fixed parameters

Name (Documentation)	Name (Programs)	Value	Description
N_s	Ns	32	number of elements in each axis
N_d	N	1024	number of design variables
N_t	Nt	6144	total number of quadrilateral elements
L_x	Lx	≈ 0.310 m	design domain shorter side
L_y	Ly	≈ 0.537 m	design domain longer side
e_x	Lex	≈ 0.00969 m	shorter side length of the elements
e_y	Ley	≈ 0.01679 m	longer side length of the elements
maximal decrease of \hat{E}	Eyvar	0.05 Pa	maximal decrease in Young’s modulus per iteration
D_{\max}	Dmax	1.5625%	maximal topology variation
r_s	rsen	0.024 m	sensitivity filter radius
r_m	rmor	0.018 m	morphology filter radius
P	patience	30	patience stopping criterion
momentum	momentum	25%	momentum value for the objective function sensitivity
β	beta	0.05	volume penalization factor
\tilde{E}	Ey	1.0 Pa	Young’s modulus of the base material
$\tilde{\nu}$	nu	0.3	Poisson’s ratio of the base material
p_k	pk	1×10^{-9}	soft-kill parameter
noptf	noptf	7	number of optimizations stored in the same file

The range of values for each input parameter is presented in [Table 2](#). The target Poisson’s ratio ranges from -1.0 to 1.0 , in steps of 0.01 . Values between 0.20 and 0.40 are not considered since they are close to the property of the base material, so the values $\{0.21, 0.22, \dots, 0.39\}$ are taken out of the set. This results in 182 possible values for ν^* . The minimal Young’s modulus ranges from 0.0% to 50.0% , in steps of 0.5% . This results in 101 possible values for E_{\min} . Therefore, there are $18382 = 182 \times 101$ unique pairs (ν^*, E_{\min}) .

Table 2: Input parameters

Name	Range of values	Description
ν^*	$-1.00 \sim 0.20 \mid 0.40 \sim 1.00$	target Poisson’s ratio
E_{\min}	$0.0\% \sim 50.0\%$	minimal Young’s modulus

The github repository (<https://github.com/Joquempo/Metamaterial-Dataset>) is structured as presented in [Figure 22](#). The “sample” folder contains scripts for taking samples from the generated dataset. The “source” folder contains the scripts that generate the dataset. The “validation” folder contains scripts used to validate the presented implementation. The “CITEAS” file lists indicated references for citing this work. The “LICENSE” file presents the terms of the GNU General Public License. The “README.md” file contain some key points from this documentation. And the “documentation.pdf” file corresponds to this document itself.

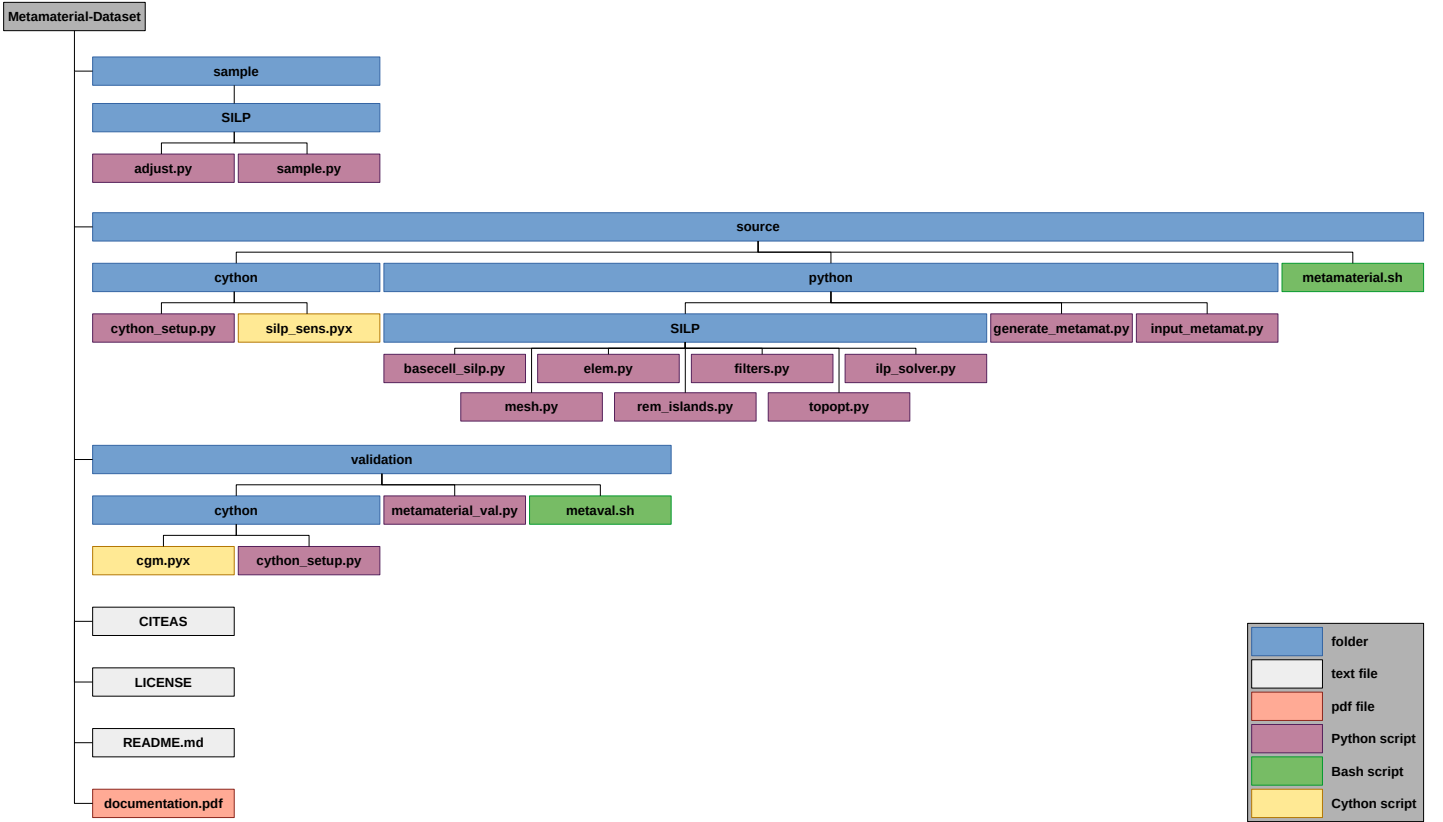


Figure 22: Metamaterial-Dataset repository

The “./source/python/SILP/basecell_silp.py” script should be edited before executed, in order to set appropriate values for the parameters “fid_ini” and “fid_lim”. For example, if “fid_ini=0” and “fid_lim=700”, the program will run the first 700 cases, $\{0, 1, \dots, 699\}$; if “fid_ini=700” and “fid_lim=2800”, it will run the subsequent 2100 cases, $\{700, 701, \dots, 2799\}$.

After installing and activating Anaconda, the dataset can be generated by executing the scripts ordered in the execution tree shown in Figure 23. The “Do” column specifies what is done by each script; the “Use” column specifies data and other scripts used during execution; the “Generate” column specifies what each script generates after being executed.

After generating the dataset, samples can be taken by using the scripts from the “sample” folder. To select which samples should be taken, the “./sample/SILP/sample.py” script must be edited. The indices of the files must be set, as well as the flags used to select what should be plotted. Subfolders will be created to store the generated figures.

Independently, the validation routine can be executed by using the scripts from the “validation” folder. The bash script “./validation/metaval.sh” can be executed to build the required cython codes. Then, after activating the “metamaterial” conda environment, the “./validation/metamaterial_val.py” script can be executed to perform the procedure. Results will be printed to the terminal. An IDE is recommended in order to visualize the validation figures (otherwise, the script should be altered so the figures are not immediately closed after concluding the validation procedure).

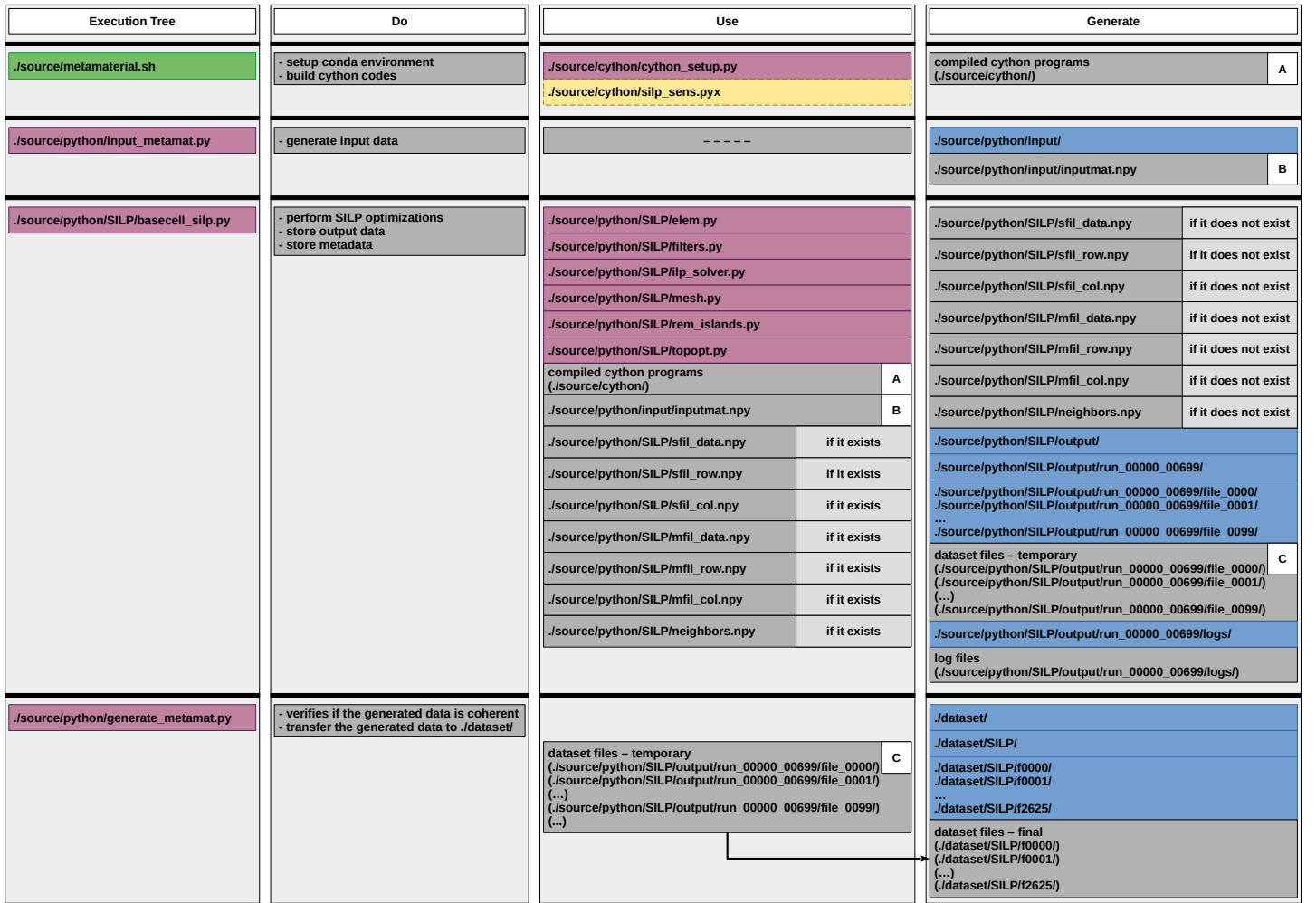


Figure 23: Execution tree

Table 3 presents the dataset files. The size shown in the second column corresponds to the amount of the useful data, considering the average value of 74 iterations for the optimization procedure. The size shown in the third column corresponds to minimal and maximal observed values for the disk usage. Each of these files contains data from 7 optimization processes, so 2626 of each one is generated after running all the 18382 cases.

Table 4 presents the disk usage for the complete dataset.

Since all data is stored as numpy arrays, through “`numpy.save()`”, everything can be easily read through “`numpy.load()`”. In anyway, the sampling script illustrates how to read data from the generated dataset. As a final remark, it should be noted that careless sampling can quickly write a lot of data in your disk, so be cautious when running the sampling script.

Table 3: Dataset files

Name	Size (useful data)	Size (observed disk usage)	Description
dC00.0.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-0 finite variations of C_{00}
dC11.0.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-0 finite variations of C_{11}
dC22.0.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-0 finite variations of C_{22}
dC00.1.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-1 finite variations of C_{00}
dC11.1.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-1 finite variations of C_{11}
dC22.1.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-1 finite variations of C_{22}
dC00.2.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-2 finite variations of C_{00}
dC11.2.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-2 finite variations of C_{11}
dC22.2.npy	2.1 MB	1.3 ~ 4.6 MB	CGS-2 finite variations of C_{22}
dC00.w.npy	2.1 MB	1.3 ~ 4.6 MB	exact finite variations of C_{00}
dC11.w.npy	2.1 MB	1.3 ~ 4.6 MB	exact finite variations of C_{11}
dC22.w.npy	2.1 MB	1.3 ~ 4.6 MB	exact finite variations of C_{22}
dis_xx.npy	25.4 MB	15.9 ~ 56.3 MB	displacements vectors used to compute C_{00}
dis_yy.npy	25.4 MB	15.9 ~ 56.3 MB	displacements vectors used to compute C_{11}
dis_xy.npy	25.4 MB	15.9 ~ 56.3 MB	displacements vectors used to compute C_{22}
Ey.npy	2.1 kB	4.1 ~ 8.2 kB	Young's modulus values
Ey_opt.npy	28 bytes	4.1 ~ 4.1 kB	optimized Young's modulus values
fid.npy	28 bytes	4.1 ~ 4.1 kB	input files indices
inp.npy	56 bytes	4.1 ~ 4.1 kB	input data
nu.npy	2.1 kB	4.1 ~ 8.2 kB	Poisson's ratio values
nu_opt.npy	28 bytes	4.1 ~ 4.1 kB	optimized Poisson's ratio values
ptr2inp.npy	2.1 kB	4.1 ~ 8.2 kB	pointers from iterations to inputs
ptr2opt.npy	32 bytes	4.1 ~ 4.1 kB	pointers from inputs to iterations
tim.npy	196 bytes	4.1 ~ 4.1 kB	execution times and number of iterations
top.npy	65.6 kB	41.0 ~ 143.4 kB	topology vectors
top_opt.npy	896 bytes	4.1 ~ 4.1 kB	optimized topologies
vol.npy	2.1 kB	4.1 ~ 8.2 kB	relative volume values

Table 4: Disk usage

	Useful data	Observed disk usage
Complete dataset	260 GB	277 GB

6 Acknowledgements

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