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Higher-order multi-resolution topology optimization using the finite cell method

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SUMMARY
This article presents a detailed study on the potential and limitations of performing higher-order multi-resolution topology optimization with the finite cell method. To circumvent stiffness overestimation in high-contrast topologies, a length-scale is applied on the solution using filter methods. The relations between stiffness overestimation, the analysis system, and the applied length-scale are examined, while a high-resolution topology is maintained. The computational cost associated with nested topology optimization is reduced significantly compared with the use of first-order finite elements. This reduction is caused by exploiting the decoupling of density and analysis mesh, and by condensing the higher-order modes out of the stiffness matrix. Copyright © 2016 John Wiley & Sons, Ltd.

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1. INTRODUCTION

In the past decades, density-based topology optimization has become a mature design method, with applications in a variety of industries. Despite the rapid advancements in computer performance, large-scale topology optimization still comes at a high-computational cost, dominated by the finite element (FE) analysis [1]. In this article, we extensively describe the advantages and limitations of multi-resolution methods to reduce this computational cost. Furthermore, we present an efficient multi-resolution topology optimization algorithm, while maintaining a high-resolution topology.

Currently, the far majority of topology optimization methods uses the same mesh for both density description and analysis. The density elements are then directly mapped on first-order FE, which due to their uniform size and shape allow for efficient assembly of the stiffness matrix. The introduction of the finite cell method (FCM) by Parvizian, Düster, and Rank showed that a decoupling of the density-mesh and analysis-mesh, in combination with higher-order shape functions, can outperform the aforementioned approach, for sufficiently smooth density distributions [2–4]. In this fictitious domain method, multiple density elements are mapped on analysis cells operating at a higher-order basis. This allows for accurate and efficient analysis of data directly derived from X-ray scans or quantitative CT scans without the need for meshing [4–7]; however, the method is also very interesting for topology optimization applications. To demonstrate this, the developers of the FCM implemented a heuristic optimization method, showing promising results [8]. In a similar approach, Nguyen et al. report a reduction in computational cost when decoupled meshes, linear shape
functions, and a gradient-based optimization method are used [9, 10]. A similar approach is applied in a very popular topology optimization app for hand-held devices [11], while in a more recent study, Nguyen et al. have demonstrated the use of higher-order shape functions in combination with this multi-resolution approach [12].

In this paper, we go a step further and dedicate a large part of our attention to the limits at which topology optimization using higher-order multi-resolution methods can be performed, because this will greatly help the method's maturation. It is known that the FCM shows superior convergence compared with first-order FE for smooth structures [13]; however, in topology optimization, highly inhomogeneous topologies belong to the solution space [14]. Filter methods are employed to impose a length-scale on the solution, and we demonstrate that the quality of the corresponding solution depends on both filter and properties of the analysis mesh. Using a large number of numerical examples for typical minimum compliance and minimum displacement problems, we find an indication of the parameters for which topology optimization using the FCM results in satisfying topologies (i.e., topologies similar to the ones obtained using standard linear FE).

All experiments shown in this paper have been performed in a MATLAB framework that is created on top of FCMLab: A Finite Cell Research Toolbox for MATLAB, developed by Zander et al. [15]. The developed framework is similar to the efficient 88-line topology optimization code [16], and the MATLAB implementation of the method of moving asymptotes (MMA) is used to solve the optimization problem [17]. Using this efficient optimization framework, we present a detailed study on the computational cost of the method and show its competitiveness compared with the use of linear FE. To do so, we present a modification to the FCM, where we condense the internal modes out of the stiffness matrix using the Schur-complement, and show a significant gain in efficiency when higher-order multi-resolution topology optimization is performed.

The paper is organized as follows: The methodology of the FCM is introduced in Section 2. In Section 3, the theory used for the topology optimization is described. The limitations of the method are identified and shown in Section 4. The corresponding tests on the efficiency of the developed method are shown in Section 5, which also includes a detailed discussion on the performance of the method. Finally, Section 6 will present the most important conclusions of this study.

2. THE VOXEL-VERSION OF THE FINITE CELL METHOD

In the voxel-version of the FCM, separate meshes are used to describe the geometry and to perform the analysis. The elements involved with the analysis mesh are called cells, while the geometry (topology) is described by density elements called voxels (volume pixels). Both cells and voxels have a uniform shape throughout the design domain and are square in 2D problems and cubic in 3D problems, as can be seen in Figure 1. The distribution of voxels within a cell can be parameterized by the amount of voxels in one cell direction ($n_{voxel}$); hence, the total amount of voxels in a cell ($n_{sc}$) depends on $n_{voxel}$ and the dimension of the design domain.

![Figure 1. The different types of meshes used in the finite cell method, with $n_{voxel} = 5$.](image)
The displacement field in a cell with a complex material distribution cannot be interpolated with sufficient accuracy using linear shape functions; therefore, the FCM includes the $p$-version of the FEM [18]. The voxel contributions to the cell stiffness matrix ($k_c$) are applied using a composed integrations scheme, that is, the stiffness matrix and load vector are integrated in the voxels and then mapped on the cells [2, 4], where the voxel stiffness is interpolated using the solid isotropic material with penalization (SIMP) method [19]:

$$k_c = \sum_{i=1}^{n_{sc}} \left( E_{\text{min}} + \tilde{\rho}_i (E - E_{\text{min}}) \right) k_i^0$$

where $\tilde{\rho}_i$ is the physical density associated with the $i$th voxel, $q$ is the penalization factor, $E$ is the Young’s modulus of a solid voxel, $E_{\text{min}}$ is a very small value ($\sim E \cdot 10^{-9}$) to avoid ill-conditioning of the stiffness matrix, and $k_i^0$ corresponds to the contribution of the $i$th voxel using a unit stiffness.

### 2.1. Higher-order shape functions

Integrated Legendre polynomials are used to form the higher-order basis. Contrary to Lagrange polynomials, Legendre polynomials are hierarchic, that is, the shape functions for polynomial degree $p$ are included in the approximation space when degree $p + 1$ is used, as can be seen in Figure 2.

The corresponding one-dimensional set of shape functions can be defined as follows:

$$N_1(\xi) = \frac{1}{2} (1 - \xi)$$

$$N_2(\xi) = \frac{1}{2} (1 + \xi)$$

$$N_i(\xi) = \phi_{i-1}(\xi), \quad i = 3, 4, \ldots, p + 1$$

where $N_i(\xi)$ corresponds to the $i$th shape function, and where $\phi$ corresponds to an integrated Legendre polynomial.

With the integrated Legendre polynomials as basis functions, the displacement field can be interpolated:

$$u(\xi) = N_1(\xi)u_1 + N_2(\xi)u_2 + \sum_{i=3}^{p+1} N_i(\xi)u_i$$

Here, $u_1$ and $u_2$ correspond to the nodal displacements, while $u_i$ corresponds to the amplitudes of the higher-order shape functions.

The one-dimensional shape functions can be used in two-dimensional or three-dimensional problems, by combining the bases in the tensor product space [15]:

$$N_{i,j}^{1D}(\xi, \eta) = N_i^{1D}(\xi)N_j^{1D}(\eta), \quad i, j = 1, 2, \ldots, p + 1$$

$$N_{i,j,k}^{2D}(\xi, \eta, \zeta) = N_{i,j}^{1D}(\xi, \eta)N_k^{1D}(\zeta), \quad i, j = 1, 2, \ldots, p + 1$$

(a) Lagrange polynomials

(b) Integrated Legendre polynomials

Figure 2. Set of one-dimensional standard (left) and hierarchic (right) shape function for $p = 1, 2, 3$ [20].
In the current code, the full tensor product space was used, due to its support in FCMLab [15]. Alternatively, the trunk space, which is a deflated version of the tensor product space, will produce an equivalent solution quality with less degrees of freedom, especially for higher-order polynomial degrees [18, 20]. On a standard quadrilateral element, three different types of modes can be distinguished, shown in Figure 3. The nodal modes and edge modes are shared with adjacent cells, while the internal modes are local to one cell.

2.2. Static condensation

A disadvantage of a higher-order basis is the large amount of internal modes with increasing p-degree, that is, modes that are specific to only one cell. Figure 4 shows the number of internal modes and the total number of modes at different p, for both 2D and 3D problems.

These internal modes can be eliminated from the global system of equations by condensing them out of the stiffness matrix, decreasing the computational cost of the analysis. Furthermore, the procedure results in a drastic decrease in the condition number, which is highly beneficial when iterative solvers are considered [21]. The global system of equations can be re-ordered such that the condensed matrix becomes the Schur complement of \( K_{ii} \) in \( K \).

\[
\begin{bmatrix}
K_{ee} & K_{ei} \\
K_{ei}^T & K_{ii}
\end{bmatrix}
\begin{bmatrix}
U_e \\
U_i
\end{bmatrix}
= \begin{bmatrix}
F_e \\
F_i
\end{bmatrix}
\quad (5)
\]

where \( U \) is the displacement vector, \( F \) is the force vector, subscript \( e \) denotes the external modes, while subscript \( i \) corresponds with the internal modes. From the second row of the system of equations, it follows that

\[
U_i = K_{ii}^{-1} \left[ F_i - K_{ei}^T U_e \right] 
\quad (6)
\]

Substitution in the first row of Equation (5) yields

\[
\begin{bmatrix}
K_{ee} - K_{ei} K_{ii}^{-1} K_{ei}^T
\end{bmatrix}
U_e = F_e - K_{ei} K_{ii}^{-1} F_i
\quad (7)
\]

![Figure 3. Two-dimensional mode types [15].](image1)

![Figure 4. Number of internal degrees of freedom (\( n_{int} \)) and external degrees of freedom (\( n_{ext} \)) in \( k_e \) as a function of \( p \) for both 2D and 3D problems.](image2)
which is the condensed system of equations,

$$ K^* U_e = F^* $$

where $U_e$ and $F^*$ denote the condensed systems displacements and loads, respectively. Because the indices of $K_{ii}$ are purely local to one cell, we can assemble the condensed stiffness matrix ($K^*$) efficiently by mapping the contributions of the condensed cell stiffness matrices ($k^*_c$):

$$ k^*_c = [k^*_{c,ee} - k^*_{c,ei} k^{-1}_{c,ii} k^T_{c,ei}] $$

3. TOPOLOGY OPTIMIZATION

Topology optimization can be seen as a material distribution problem. The goal is to find an optimum material distribution that minimizes an objective function $O_F$. This function is subject to $m$ constraints $O_G$, of which the first is generally a volume constraint. The design domain $\Omega$ is discretized into voxels (volume pixels) to which design variables are assigned, all variables together form the design vector $\rho$. The material distribution is allowed to vary between 0 and 1 for gradient-based optimization.

The discretized topology optimization problem will have a mesh-dependent solution. Furthermore, numerical artifacts, similar to the well-known checkerboard patterns, need to be omitted from the solution space. To do so, several established filter methods are used in our model, such as the sensitivity filter, the density filter, the density filter with projection, and robust topology optimization [22]. The description of each of these filters and their corresponding sensitivities can be found in Appendix A. Using these filter methods, the design vector $\rho$ is linked to the physical density in each voxel $\tilde{\rho}$; hence, the discretized optimization problem can be written as [23]:

$$ \min_{\rho} : \tilde{F}(\rho) = F(\rho, U_e) $$

$$ \text{s.t. : } K^* U_e = F^* $$

$$ : \tilde{G}_1(\rho) = v^T \tilde{\rho} - V_{max} \leq 0 $$

$$ : \tilde{G}_i(\rho) = G_i(\rho, U_e) \leq 0, \quad i = 2, \ldots, m $$

$$ : 0 \leq \rho \leq 1 $$

where $v$ is the vector containing the element volumes, and $V_{max}$ is the maximum allowed volume of the material in the design domain. The optimization problem described earlier is a nested topology optimization problem, that is, the equilibrium equations are satisfied for each optimization step using the FCM. For the design update, the MATLAB implementation of the MMA is used [17].

3.1. Definition of test-problems

In this study, four representative test-problems are used to test the limitations and the computational cost of performing topology optimization with the FCM. The MBB-beam, which is a typical benchmark example, is used to test the behavior for minimum compliance problems, where, because of the symmetry of the design problem, we model only half of the beam. In another, more challenging minimum compliance problem, a cantilever beam, is subject to a uniform traction load. This will demonstrate the performance of the method in regions subject to very small loads. Furthermore, the well-known compliant force inverter is used to identify the performance of the method for mechanism design problems, and the ability of the method to form hinges [24]. Finally, a 3D, and slightly shortened version of the MBB-beam is used to test the computational cost of topology optimization with the FCM. Sketches of the domain and boundary conditions of each of these examples can be found in Figure 5.

In all 2D-examples, plane stress conditions are assumed. For the 2D-version of the MBB-beam, we chose $F = 1$, for the cantilever-beam subject to a distributed load $F = 1/L$, and for both cases $V^* = 0.4$. For the 3D-version, the domain is extended to the z-direction with depth $L/2$, while the domain length in the y-direction is increased from $L/3$ to $L/2$, the corresponding maximum volume
is $V^* = 0.12$. Similar to the domain, the boundary conditions are extended in the $z$-direction, where the load is now applied as a line load $F = 1/L$. For the mechanism design, the objective is to minimize the displacement $u_{\text{out}}$ for a given input force $f_{\text{in}} = 1$. The spring coefficients used are $k_{\text{in}} = 1$, and $k_{\text{out}} = 0.001$, and $V^* = 0.3$. For all optimization problems, $L$ is a unit length, a unit Young’s modulus is used, $v = 0.3$, a penalization factor $q = 3$ is used, leaving just the polynomial degree $p$, the amount of voxels per cell direction $n_{\text{voxel}}$, and the filter radius $R$ as free parameters.

A common formulation to write the objective is

$$\min_\rho \hat{F}(\rho) = \mathbf{L}_e^T \mathbf{U}_e$$  \hspace{1cm}  (11)

where vector $\mathbf{L}_e$ takes different forms for the different types of problems. In the minimum compliance problems, $\mathbf{L}_e = \mathbf{F}^*$, while for the mechanism design problem, $\mathbf{L}_e$ is a vector which contains all zeros except for the index corresponding to $u_{\text{out}}$, which is set to one. The sensitivity of the objective w.r.t. physical density $\tilde{\rho}_i$ can be calculated by adjoint sensitivity analysis [25],

$$\frac{\partial \hat{F}}{\partial \tilde{\rho}_i} = \lambda_{c,e}^T \left( \frac{\partial k_{c,ee}}{\partial \tilde{\rho}_i} - \frac{\partial k_{c,ei}}{\partial \tilde{\rho}_i} k_{c,ii}^{\text{inv}} k_{c,ei}^{\text{inv}} \right) + \lambda_{c,ei} \left( k_{c,ii}^{\text{inv}} \frac{\partial k_{c,ei}}{\partial \tilde{\rho}_i} - k_{c,ei} k_{c,ii}^{\text{inv}} \frac{\partial k_{c,ei}}{\partial \tilde{\rho}_i} - k_{c,ei} k_{c,ii}^{\text{inv}} \frac{\partial k_{c,ei}}{\partial \tilde{\rho}_i} \right) \mathbf{u}_{c,e}$$  \hspace{1cm}  (12)

where $\lambda$ is the adjoint vector that can be obtained using

$$\lambda_{c,e}^T = -(\mathbf{K}^*)^{-1} \mathbf{L}_e^T$$  \hspace{1cm}  (13)

it can be seen that for compliance minimization problems $\lambda = -\mathbf{U}_e$.

4. LIMITATIONS OF HIGHER-ORDER MULTI-RESOLUTION TOPOLOGY OPTIMIZATION

The solution space in topology optimization examples consists of highly inhomogeneous topologies. When first-order FE are utilized, these inhomogeneities can be exploited by the optimizer, resulting
in checkerboard patterns, where the stiffness of these checkerboards is overestimated [14, 26]. To circumvent this problem, filter methods are used to impose a length-scale on both material and void. In a similar fashion, the heterogeneities in the solution space can be utilized when higher-order multi-resolution topology optimization is performed. To demonstrate this, consider the MBB-beam optimization example, where $p = 4$, and $n_{\text{voxel}} = 5$. In Figure 6(a), a solution is shown, where no filter method is used.

### 4.1. The need for restriction of the solution space

The compliance $c_{\text{obj}}$ of the highly heterogeneous solution in Figure 6(a) cannot be estimated correctly by the analysis model ($p = 4$, $n_{\text{voxel}} = 5$). The solution space of the displacement field does not allow the sharp variations in strains or displacement corresponding to these heterogeneous cells; hence, the displacement is underestimated, making the cells artificially stiff. When the compliance is recalculated in a post-verification step $c_{\text{post}}$, using a highly accurate analysis model ($p = 3$, $n_{\text{voxel}} = 1$), it can be seen that the stiffness of the structure is greatly overestimated.

This problem, where the solution field cannot be computed accurately over highly non-smooth regions, is well known, and generally resolved using a local overlay mesh [4]. For topology optimization applications, the topology is not known a-priori, and hence, we argue that an imposed length-scale on both solid and void will resolve the problem as well. Here, the length-scale is imposed using the mesh-independent filter methods (Appendix A) that regularize the solution. If the filter fully covers an analysis cell, the occurrence of artificially stiff patterns is banned from the solution space. Nevertheless, we argue that the introduction of a length-scale also works well when the filter is much smaller than the analysis cell if $p$ is high enough. The introduction of this small length-scale will make sure that the presence of a density discontinuity within a cell can be captured by a sufficiently accurate analysis model.

In topology optimization problems, the goal is to locally maximize the mutual energy term $\lambda_{c,e}^T k_{c,e}^* u_{c,e}$ (equal to the strain energy density for minimum compliance problems), such that the objective is minimized. Therefore, to obtain well-connected structures, it is sufficient that the value of $\lambda_{c,e}^T k_{c,e}^* u_{c,e}$ is worse for discontinuous structures compared with the value for well-connected designs and that this negative effect is sufficiently reflected in the design sensitivities. If this is the case, the solution will be directed towards a well-connected design; hence, a very accurate representation of the solution field over the discontinuity is not required.

To demonstrate this proposition, we go back to the MBB-beam problem of Figure 6(a) where the strain energy density of the discontinuous cells is greatly overestimated. In Figure 6(b), it is
shown that an imposed length-scale, although even small, already results in an acceptable topology that is exactly similar to the design obtained when first-order FE is used, shown in Figure 6(c). The introduction of a length-scale also works well for high \( n_{\text{voxel}} \) in combination with a high polynomial degree, as explained earlier. This is illustrated by Figure 6(d), where the cell size is 15 voxel-lengths \( h \), while the filter radius is just \( 2h \).

A relation exists between the filter method on the one side and the quality of the analysis model on the other side, determining whether an acceptable solution can be obtained. An analysis model of low quality, that is, low \( p \), high \( n_{\text{voxel}} \), combined with a large \( R \) may result in an acceptable solution; however, a high-quality analysis model with small filter radius \( R \) may still overestimate the cell stiffness in the presence of a density discontinuity. In the previous examples, the chosen filter-radius \( R = L/30 \) corresponds to two voxel-lengths \( h \). In the following, we specify \( R \) in terms of \( h \), because this is an indicator for the locally imposed length-scale on the physical density and therefore directly linked to the occurrence of these artificially stiff patterns.

4.2. The validity of optimized solutions

The solutions obtained using the FCM as analysis model have to be compared with the well-established solutions obtained using linear FE. When these solutions are similar in both performance and topology, they are deemed acceptable/satisfactory.

To test the performance of the optimized designs, we propose two different methods to check the validity of the results:

- **Post-verification**: It is extremely important to post-verify the objective and constraints, with a high-quality analysis model \((p = 3, \ n_{\text{voxel}} = 1)\). The artificially stiff patterns shown in Figure 6(a) can be immediately identified by comparing \( c_{\text{post}} \) and \( c_{\text{obj}} \). Furthermore, post-verification of the objective is the only way to obtain a fair comparison between generated topologies using different analysis models.

- **Visual check**: The effect of a discontinuity in a low-load region does not always translate into a large difference between \( c_{\text{post}} \) and \( c_{\text{obj}} \). For the problem where the cantilever is subject to a distributed load, disconnected patches of material may occur in the low-load region of the upper-right corner, as can be seen in Figure 7. The effect of these disconnected regions on the behavior of the structure is small; hence, they should be identified via a visual check of the solution.

The visual check may not always work for large 3D examples, for example, discontinuities can exist within a closed cell. However, in this case, one can also consider the strain energy density for the solid voxels in the post-verified solution. If these strain energy densities are several orders of magnitude smaller than for other solid voxels, these solid voxels are non-load carrying and indicate the presence of a discontinuity.

![Figure 7. Optimization of the distributed load optimization example using a discretization of 120×60 voxels, using sensitivity filtering with \( R = 2h \).](image-url)
4.3. The occurrence of artificially stiff patterns in low-load regions

Artificially stiff patterns are most likely to occur in regions where the sensitivity analysis cannot sufficiently reflect the difference between a well-connected and an artificially stiff pattern. This effect can be captured best by the the distributed load problem (Figure 7), where at the low-load regions the strain energy density is small compared with the rest of the domain. Correspondingly, the sensitivities are very small; hence, it can be more beneficial to have a disconnected structure with a slightly lower strain energy density than a well-connected structure that requires more material. Therefore, the analysis model has to have a sufficiently high $p$, such that the negative effect of an artificially stiff pattern can be reflected in the design sensitivities, and hence, the optimizer will end up with a well-connected design.

If a sensitivity or density filter is used, it is also possible that gray material, that is, material of intermediate density, is introduced at these low-load regions. This happens as well when first-order FE are used, for example, Figure 8(a). However, if these regions of gray material show some local variation between dense and less-dense voxels, as is the case in Figure 7(a) and (b), then this is still considered an unacceptable solution. This local variation between dense and less-dense voxels is caused by the optimizer exploiting the analysis model and will never occur when first-order FE are used. Therefore, even small variations as in Figure 8(b) are deemed unacceptable.

4.4. The effect of the filter method and filter radius

Maximum design resolution is obtained for a small filter radius $R$. However, a small $R$ requires a very accurate analysis model, which means that the method will be computationally much more expensive than the use of first-order FEs. Therefore, the smallest value of $R$ has to be found, which allows well-connected solutions for an analysis system of moderate quality. To do so, all 2D experiments have been performed using sensitivity filtering, for different $R$, $p$, for a fixed $n_{\text{voxel}} = 10$. We have chosen to keep a fixed, large number of voxels per cell to allow for structural members smaller than the cell-size. The lowest polynomial degrees for each $R$ that do not result in discontinuous structures can be found in Figure 9.

It can be seen that the cantilever beam subject to a distributed load is by far the most critical example, because of the low load or density regions. The MBB-beam and the compliant force inverter example show a similar but less critical behavior for small filter radii. Furthermore, it can be seen that a filter-radius of $1.8h$ or larger requires an analysis system of moderate $p$-degree. We choose to use a filter radius of $2h$ in the remainder of this study.

Besides the filter radius, the different filter methods have a different effect on the occurrence of artificially stiff patterns. A detailed description of these filter methods can be found Appendix A. In Figures 8(a) and 10, the solutions obtained for the distributed load example are shown for all different filter methods using $p = 6$, and $n_{\text{voxel}} = 10$. As indicated in Figure 9, it is not advised to use these settings, and for all examples, the obtained solutions were not deemed acceptable. However, these unacceptable topologies are good indicators of the differences between the filter methods. For both sensitivity and density filter, the transition between solid and void is gradual; hence, the
Figure 9. Lowest polynomial degree $p$ that did not result in artificially stiff patterns vs filter radius $R$, for three different test-problems, using sensitivity filtering.

Figure 10. Optimization of the distributed load optimization example using $p = 6$, $n_{\text{voxel}} = 10$, and a discretization of $240 \times 120$ voxels. Different filter methods are used with $R = 2h$.

patches of material are not completely disconnected. The density filter using the Heaviside projection can easily end up with an artificially stiff pattern, because it only imposes a length-scale on the solid. Therefore, this filter method will not be considered any further in this study. The modified Heaviside projection filter normally performs better, because it imposes a length-scale on the void. However, because it does not put a length-scale on the solid, very thin structural members can occur. In this example, one of the structural members ends abruptly, as can be seen in the top right corner. The analysis model is not able to capture this disconnection, and hence, this point is artificially stiff. Finally, it can be seen in Figure 10(d) that the robust formulation imposes a length-scale on both solid and void. At first sight, the structure seems to perform well; however, a close-inspection of the low-load region reveals that the structure performs worse compared with a reference solution obtained with first-order FE. The optimizer thus ended up at an artificially stiff local minimum; hence, the solution is regarded as unacceptable.
4.5. Experiments on the limits of higher-order multi-resolution topology optimization for R=2h

To attain a good overview of the limits at which higher-order multi-resolution topology optimization can be performed, all 2D test-problems were solved for different \( p, n_{\text{voxel}} \), for two mesh sizes (i.e., coarse and fine) and for the different filter methods (sensitivity filtering, density filtering, density filtering with the modified Heaviside projection, and robust topology optimization) all with \( R = 2h \). In total, more than a thousand different experiments have been performed, of which the most important observations are summarized as follows:

- **Similarities in generated topologies:** The acceptable optimized topologies (i.e., no artificially stiff patterns) are very similar to the optimized topologies obtained when first-order FE is used. Because of the self-adjoint nature of minimum compliance problems, the resulting topologies are almost identical, as can be seen in Figure 6 (b)–(d). The compliant force inverter example is more prone to end up at local minima due to the more complex nature of the objective function. Therefore, the solutions show more variation as can be seen in Figures 11 and 12; however, their corresponding post-verified objectives are all in the same range.

- **Effect of analysis model on local minima:** The type of analysis model does not seem to have an effect on the optimizer getting stuck in strong local minima. The use of higher-order shape functions allows for hinges at cell nodes or within a cell as can be seen in Figures 11 and 12. For all the performed experiments, we could not identify that one location for a hinge was favored over another; furthermore, the hinge was able to move freely during the design iterations.
Effect of the cell size: A larger $n_{\text{voxel}}$ requires a higher polynomial degree to prevent the formation of discontinuities within a cell. All experiments have shown that the FCM works very well as analysis model up to $n_{\text{voxel}} = 15$ or even $n_{\text{voxel}} = 20$, see for example Figures 6(d) and 12. However, a large value of $n_{\text{voxel}}$ increases the amount of possibilities in which artificially stiff patterns can occur. Larger values of $n_{\text{voxel}}$ thus require more degrees of freedom to prevent the occurrence of these undesired patterns compared with the use of a smaller number of voxels per cell, for example, $n_{\text{voxel}} < 8$. Furthermore, the combination of a high-polynomial degree and a high number of voxels requires the storage of a large number of voxel contributions to the cell stiffness matrix $k^0_i$, especially for 3D-problems. Thus, the cell size poses a limit on the computational cost, and no values of $n_{\text{voxel}} > 15$ are recommended.

Effect of filter methods: The different filter methods have a different effect on the occurrence of artificially stiff patterns. However, we observed great similarities for the settings of the analysis method that did result in artificially stiff patterns. If a type of analysis system produced an acceptable topology using the sensitivity filter, then this analysis system was almost guaranteed to work as well with the density filter, or with the robust topology optimization formulation. In a few cases that work well for the other filter methods, the density filter with the modified Heaviside projection produces artificially stiff patterns, and this is because no length-scale is posed on the solid part as is discussed in Section 4.4. Nevertheless, this filter method worked well in almost all cases.

Effect of the mesh-size: All 2D experiments have been performed on two different mesh sizes to find the effect of the mesh-size. The MBB-beam has been modelled on a mesh of $180 \times 60$ elements, and on a mesh of $360 \times 180$ elements. The cantilever beam is modelled on a coarse mesh of $120 \times 60$ elements, and on a fine mesh of $240 \times 120$ elements, while for the force-inverter example, a mesh of $120 \times 60$ and a mesh of $160 \times 80$ has been used. No differences could be found between the experiments performed on a coarse mesh and the experiments performed on a fine mesh. The reason is that the filter radius has been linked to the voxel-width $h$.

Artificially stiff patterns in the first iterations: Artificially stiff patterns can arise during the first 10–100 iterations, as can be seen in Figure 13. Here, the stiffness of the horizontal structural member, which is going through the middle of the top cells, is overestimated. Normally, these patterns are gradually removed by the filter methods, yielding an acceptable solution. This effect can still be undesired when topology optimization is performed interactively [11]. Furthermore, it is not always the case that the filter methods are able to remove these patterns, recall the topology obtained using the robust formulation in Figure 10(d), where this exact same structural member is part of the final design. This effect is more likely to occur when a cell consists of a large number of voxels, hence too large values for $n_{\text{voxel}}$ should be avoided.

Artificially stiff patterns in 3D: Visualization of 3D topologies can be carried out with a data visualization program, for example, ParaView. Similar to artificially stiff cells in 2D, the artificially stiff cells in a 3D problem consist of disconnected regions of material, and it is interesting to note that they seem to occur at the exact same settings for the analysis system as

![Figure 13. Optimization of the distributed load optimization example using $p = 7$, $n_{\text{voxel}} = 10$, and a discretization of $120 \times 60$ voxels. Sensitivity filtering is applied with $R = 2h$, $c_{obj} = 14.9$ J, and $c_{post} = 15.5$ J.](image)
in the 2D MBB-beam example. In Figure 14, the difference between an acceptable topology (a), and one with artificially stiff patterns (b) can be seen clearly. The values for \( c_{post} \) were obtained with \( p = 2 \), and \( n_{voxel} = 2 \), to avoid memory problems in MATLAB.

- **Effect of the optimization problem:** Because of the occurrence of artificially stiff cells in low-load regions, the optimization problem with the distributed load is more challenging and does not work for all combinations of \( p \) and \( n_{voxel} \) that work for the MBB-beam and the force inverter problems. This shows that the choice of analysis model depends on the optimization problem and corresponding boundary conditions. Furthermore, this shows that the distributed load optimization problem is good for finding the limits of a multi-resolution analysis method and should be considered when a new method is tested.

All experiments showed great similarities in the settings for the analysis model that prevented the occurrence of artificially stiff patterns. Therefore, these settings can be summarized in Table I. The dark gray colored cells correspond to settings which in some or all of the tested experiments resulted in artificially stiff patterns. For the light-gray cells, the occurrence of artificially stiff patterns depends on the type of optimization problem and filter method. In some cases, these settings resulted in artificially stiff patterns for the distributed load problem, but not for the other optimization problems. Furthermore, the density filter using modified Heaviside projection is not guaranteed to work. Therefore, these settings should be used with caution, and post-verification of the results is advised to make sure the optimized solution is correct. The plain cells indicate settings yielding acceptable solutions, and based on our numerical experiments, we believe that these settings are representative for the settings that can be safely used in comparable topology optimization problems. However, it is important to note that the presented results are not fully conclusive and should be interpreted as an indication of which settings are prone to these artificially stiff patterns. The quality of a solution depends on numerous parameters, such as, the type of optimization problem, boundary conditions, and filter method, and settings indicated here as acceptable might in some cases still result in overestimation of the stiffness. We can conclude that higher-order multi-resolution topology optimization can work for a large number of different analysis systems. These findings can be summarized as a (conservative) rule of thumb that holds for experiments performed with a filter radius of \( R = 2h \),

\[
p \geq \text{round}(0.75 \, n_{voxel})
\]  

Furthermore, we observed that for \( n_{voxel} \geq 8 \) there are settings for which we cannot confidently say that they will not result in artificially stiff patterns. In a future study, it would be interesting to reduce the amount of light-gray cells by looking into the use of a local overlay mesh at high-contrast regions [4]. Finally, in terms of accuracy, the settings that are indicated by the plain cells
showed numerical convergence properties that are in line with the accuracy expected of the FCM method [4]. The normalized error between $c_{post}$ and $c_{obj}$ showed that $p$-refinement is more beneficial than $h$-refinement.

Table I. Results of the experiments on the limitations of higher-order multi-resolution topology optimization.

<table>
<thead>
<tr>
<th>$n_{\text{voxel}}$</th>
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<tr>
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</tbody>
</table>

The dark gray colored cells indicate settings that yield artificially stiff patterns, while for the light-gray settings, the occurrence of artificially stiff patterns depends on the type of problem and filter method. The settings for the plain cells are likely to result in acceptable topologies.

5. NUMERICAL EXPERIMENTS ON THE EFFICIENCY OF THE METHOD

The numerical cost of performing topology optimization is dictated by the cost of the repeated solving of the analysis equations. In the framework used for this study, the solution is obtained using the direct solvers implemented in MATLAB. For sparse matrices, the amount of operations is $O(n_e^{3/2})$ for 2D problems, and $O(n_e^3)$ for 3D problems, where $n_e$ is the size of the condensed stiffness matrix [27]. The actual cost of the solution also depends on the order of the approximation, linear shape functions will yield a highly diagonal stiffness matrix, while a higher-order basis will increase the bandwidth, thus increasing the corresponding computational cost. With increasing $p$, the amount of internal modes $n_{c,i}$ grows exponentially. For 3D optimization examples, the inversion of $k_{c,ii}$, which costs $O(n_e^3)$ operations, can thus have a large influence on the computational cost.

To show the computational cost for different values of $p$ and $n_{\text{voxel}}$, two different optimization examples will be discussed. The cantilever beam subject to a distributed load is used with a discretization of $240 \times 120$ voxels, and the 3D version of the MBB-beam is used with a discretization of $80 \times 40 \times 40$ voxels. In both optimization examples, sensitivity filtering is applied with $R = 2h$. All optimization examples were solved using a single-core MATLAB code on a standard laptop PC. The results for the 2D optimization example can be seen in Table II. For the 3D MBB-beam, it was not possible to test all settings for $n_{\text{voxel}}$, due to problem discretization enforced by hardware limitations. The settings that could be tested are shown in Table III. Please note that the setting for the analysis model resulting in artificially stiff patterns, as well as the settings expected to be computationally more costly than first-order FE, have been disregarded, where the settings marked with an asterisk (*) indicate that they are more costly than first-order FE at comparable quality of the analysis results.

It can be seen that decoupling of the density-mesh and analysis-mesh can be computationally more efficient than topology optimization using first-order FE. For 2D problems, an increase in speed of 2.9 can be achieved while maintaining a high-resolution topology without artificially stiff patterns. For the settings where we are unsure whether artificial patterns can occur, a speed-up of
Table II. Normalized speed-up per design iteration for the cantilever beam, subject to a uniform pressure load.

<table>
<thead>
<tr>
<th>$n_{\text{voxel}}$</th>
<th>1</th>
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<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
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<tr>
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<td>0.92*</td>
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<td>3.55</td>
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<tr>
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<td>*</td>
<td>0.60*</td>
<td>1.00</td>
<td>1.48</td>
<td>1.87</td>
<td>2.82</td>
<td>3.59</td>
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<td>*</td>
<td>0.38*</td>
<td>0.69*</td>
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<td>2.87</td>
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<td>*</td>
<td>*</td>
<td>*</td>
<td>0.29*</td>
<td>0.41*</td>
<td>0.63*</td>
<td>0.88*</td>
<td>1.17</td>
<td>1.58</td>
</tr>
</tbody>
</table>

A discretization of 240 by 120 voxels is used. The settings marked with an asterisk (*) indicate that they are computationally heavier than first-order FE.

Table III. Normalized speed-up per design iteration for the 3D MBB-beam example.

<table>
<thead>
<tr>
<th>$n_{\text{voxel}}$</th>
<th>1</th>
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<td>*</td>
<td>*</td>
<td>12.9</td>
<td>31.8</td>
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<td>$p = 4$</td>
<td>*</td>
<td>*</td>
<td>1.68</td>
<td>7.81</td>
<td>46.4</td>
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</tr>
<tr>
<td>$p = 5$</td>
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<td>*</td>
<td>*</td>
<td>0.83*</td>
<td>18.4</td>
<td>35.0</td>
</tr>
<tr>
<td>$p = 6$</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>4.13</td>
<td>14.6</td>
</tr>
</tbody>
</table>

A discretization of 80 by 40 by 40 voxels is used. The settings marked with an asterisk (*) indicate that they are computationally heavier than first-order FE.

3.6 can be achieved. Because this is a minor difference, we would recommend to perform topology optimization with settings $(p = 2, n_{\text{voxel}} = 3), (p = 3, n_{\text{voxel}} = 5), (p = 4, n_{\text{voxel}} = 6), (p = 6, n_{\text{voxel}} = 8)$ or $(p = 7, n_{\text{voxel}} = 10)$. A higher value for $p$ or $n_{\text{voxel}}$ shows a slight increase in computational cost due to the inversion of $k_{c,ii}$.

For 3D problems, the computational cost can be reduced even more. In this optimization example, no artificially stiff patterns have been spotted for the settings in the light-gray cells, as can be seen in Figure 14(a); hence, a speed-up in computational time of a factor 67 has been achieved. Still, we would not recommend to use these settings blindly, especially because the more ‘robust’ analysis model $(p = 3, n_{\text{voxel}} = 5)$ results in a computational speed-up of 32. It is interesting to note that in the 3D-examples a moderately high $p$ seems to be best. The use of $p > 4$ drastically increases the cost of the inversion of $k_{c,ii}$; furthermore, a higher $p$ increases the number of non-zero elements in $K^*$, which put a large burden on the memory requirements.

Optimization of the 128,000 design variables using first-order FE costs around 300 seconds per design iteration. When we use $p = 3$, and $n_{\text{voxel}} = 5$, this is reduced to 9.5 seconds per iteration, resulting in a total optimization time of only 34 min. We are confident that the optimization time can be reduced even further by using an efficient multigrid pre-conditioned iterative solver [28]. The challenge here will lie in finding an efficient multigrid algorithm suited for higher-order methods, which cannot be exploited by the artificially stiff patterns.

6. CONCLUSION

An efficient approach to perform higher-order multi-resolution topology optimization using voxel-version of the FCM was presented. The most important finding is that a multi-resolution analysis
The model can overestimate the stiffness of highly inhomogeneous patterns, in a similar fashion as first-order FE overestimate the stiffness of checkerboard patterns. This problem can be resolved using mesh-independent filter techniques, where we showed the relation between the quality of the analysis model and the imposed length-scale. Using more than thousand examples, we demonstrated the limits at which topology optimization, in the framework of higher-order multi-resolution methods, can be performed. Based on a number of representative test-cases, we have identified settings for the analysis models for which acceptable solutions were achieved.

By reducing the size of the stiffness matrix using static condensation, we demonstrated that the computational cost can be decreased significantly compared with the use of first-order FE. In 2D optimization examples, an increase of speed of a factor 2.9 was achieved, while in 3D topology optimization problems, an even more promising speed-up of 32 was possible. A 3D topology optimization problem with 128,000 design elements was optimized on a standard PC in 34 min, using the direct solvers implemented in MATLAB. This overall promising performance paves the way for further development of the methodology by using efficient multigrid pre-conditioned iterative solvers. We are confident that this will show further reduction in computational cost and might further reveal the potential of the method for large-scale topology optimization.

APPENDIX A: FILTER METHODS USED IN THIS STUDY

A.1. Sensitivity filtering

For the sensitivity filter introduced by Sigmund [24], no difference exists between the physical density and the design vector \( \rho = \tilde{\rho} \); however, the voxel sensitivity is now based on the sensitivities of the surrounding voxels within a mesh-independent radius \( r_{\text{min}} \). These filtered sensitivities are then used to update the design vector,

\[
\frac{\partial \hat{F}}{\partial \rho_e} = \frac{1}{\max(\rho_e, 0.001) \sum_{i=1}^{n_e} H_{ei} \rho_i} \sum_{i=1}^{n_e} H_{ei} \rho_i \frac{\partial \hat{F}}{\partial \rho_i}
\]

where \( n_e \) is the number of voxel, and \( H_{ei} \) is a linear decaying weighting function. The small number is put in the denominator to avoid division by zero. \( H_{ei} \) depends on the distance between the voxel center and the center of the surrounding voxel, as well as the filter radius \( r_{\text{min}} \):

\[
H_{ei} = r_{\text{min}} - \text{dist}(e, i)
\]

A.2. Density filtering

An alternative to the sensitivity filter is the density filter [29, 30]. The physical density of a voxel \( \tilde{\rho}_e \) is defined as the weighted average of the design variables of neighboring voxels in \( r_{\text{min}} \):

\[
\tilde{\rho}_e = \frac{1}{\sum_{i=1}^{n_e} H_{ei}} \sum_{i=1}^{n_e} H_{ei} \rho_i
\]

Using the chain rule, the sensitivities with respect to the design variables can be obtained:

\[
\frac{\partial \hat{F}}{\partial \rho_e} = \sum_{i=1}^{n_e} \frac{\partial \hat{F}}{\partial \tilde{\rho}_i} \frac{\partial \tilde{\rho}_i}{\partial \rho_e}
\]

A.3. Density filtering using a projection

A disadvantage of both sensitivity and density filter is that they introduce regions with intermediate densities. To cope with this, a projection scheme based on a smoothed Heaviside function has been introduced by Guest et al. [31]. In this approach, the filtered densities \( \tilde{\rho}_e \) using the density filter are
now referred to as the ‘intermediate design vector’. The physical density \( \tilde{\rho} \) can be calculated using the threshold method presented by [32],

\[
\tilde{\rho}_e = \frac{\tanh(\beta \eta) + \tanh(\beta(\tilde{\rho}_e - \eta))}{\tanh(\beta \eta) + \tanh(\beta(1 - \eta))}
\]  

(A.5)

where \( \eta \) is the threshold parameter, and \( \beta \) controls the smoothness of the Heaviside function. For \( \beta = 0 \), the filter gives exactly the same output as the density filter, whereas when \( \beta \) goes towards infinity, the Heaviside function is approximated. For \( \eta = 0 \), the projection corresponds to the Heaviside projection that applies a length-scale on the material, while \( \eta = 1 \) corresponds to the modified Heaviside projection introduced by Sigmund that applies a length-scale on the void regions [19]. In the experiments, an initial value of \( \beta = 1 \) is used, which is doubled every 50 iterations until a maximum of \( \beta = 64 \). The sensitivity of the objective function w.r.t. a design variable \( \rho_e \) can be written as follows:

\[
\frac{\partial \hat{F}}{\partial \rho_e} = \sum_{i=1}^{n_e} \frac{\partial \hat{F}}{\partial \tilde{\rho}_i} \frac{\partial \tilde{\rho}_i}{\partial \rho_e}
\]  

(A.6)

### A.4. Robust topology optimization

Robust topology optimization has been introduced by Sigmund as a method to perform manufacturing tolerant topology optimization [33]. Small changes in manufacturing should not lead to large changes in functionality. Another positive effect of this method is that it is able to put a length-scale on both solid and void material, eliminating the longstanding problem of one-node connected hinges [34].

In robust topology optimization, three different designs are formulated based on the same design vector. These are a dilated (\( \tilde{\rho}^d \)), intermediate (\( \tilde{\rho}^i \)), and eroded (\( \tilde{\rho}^e \)) design, with thresholds \( \eta \), 0.5, and \( 1 - \eta \), respectively. The optimization problem is now reformulated as a \( \min - \max \) problem. The sensitivities can be obtained using Equation (A.6). Analogous to [34], the volume constraint is imposed on the dilated design and updated every 20 iterations.

### ACKNOWLEDGEMENTS

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### REFERENCES


