Fast optimization-based conservative remap of scalar fields through aggregate mass transfer

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Abstract

We develop a fast, efficient and accurate optimization-based algorithm for the high-order conservative and bound-preserving remap (constrained interpolation) of a scalar conserved quantity between two close meshes with the same connectivity. The new formulation is as robust and accurate as the flux-variable flux-target optimization-based remap (FVFT-OBR) [1, 2] yet has the computational efficiency of an explicit remapper. The coupled system of linear inequality constraints, resulting from the flux form of remap, is the main efficiency bottleneck in FVFT-OBR. While conventional remappers use the flux form to directly enforce mass conservation, the optimization setting allows us to treat mass conservation as one of the constraints. To take advantage of this fact, we consider an alternative mass-variable mass-target (MVMT-OBR) formulation in which the optimization variables are the net mass updates per cell and a single linear constraint enforces the conservation of mass. In so doing we change the structure of the OBR problem from a global linear-inequality constrained QP to a singly linearly constrained QP with simple bounds. Using the structure of the MVMT-OBR problem, and the fact that in remap the old and new grids are close, we are able to develop a simple, efficient and easily parallelizable optimization algorithm for the primal MVMT-OBR QP. Numerical studies confirm that MVMT-OBR is as accurate and robust as FVFT-OBR, but has the same computational cost as the explicit, state-of-the-art FCR.

Keywords: constrained interpolation, remap, flux-corrected remap, FCT, optimization-based remap, quadratic programming

1. Introduction

We develop a fast, efficient and accurate optimization-based algorithm for the high-order conservative and bound-preserving remap (constrained interpolation) of a scalar conserved quantity between two close...
meshes with the same connectivity. This task originates in Arbitrary Lagrangian-Eulerian (ALE) methods [3], which are the main motivation for our work. In the ALE context we are given the mean value of the primitive variable (an unknown positive scalar function, such as density) on each cell of the old (Lagrangian) mesh. The conserved variable, such as mass, is the product of this mean value and the cell volume. The objective is to find an accurate approximation of the mass on the new (rezoned) mesh. The remapped cell mass divided by the volume of the new cell approximates the density, which must satisfy physically motivated bounds.

The paper continues the development and study of the optimization-based remap (OBR) approach initiated in [1, 2]. There we rephrase remap as a global inequality-constrained quadratic program (QP) for the mass fluxes exchanged between neighboring cells. The objective is to minimize the distance between these fluxes and some given target fluxes subject to constraints that enforce physically motivated local bounds on the primitive variable (density). The resulting flux-variable flux-target (FVFT) OBR has valuable theoretical and computational properties, which set it apart from advection-based [4, 5, 6], mass “repair” [7, 8], or flux-correction motivated [9] algorithms.

In a nutshell, these methods invoke local “worst-case” scenarios to preserve the local bounds through monotone reconstruction, mass redistribution, or convex combinations of low and high-order fluxes. This entangles accuracy considerations with the enforcement of the bounds, which tends to obscure the sources of discretization errors and complicates the analysis of the algorithms.

In contrast, the OBR strategy completely separates accuracy from the enforcement of the physically motivated local bounds. The latter define a feasible set for the QP, whereas the minimization of the objective function enforces the former. As a result, FVFT-OBR always finds a globally optimal, i.e., the best possible, flux-variable flux-target (FVFT) OBR solution. This allows us to exploit the structure of the problem, which can be efficiently solved using methods like the Newton or interior point methods.

Thorough convergence studies in [1] confirm that the FVFT-OBR formulation is as accurate as the state-of-the-art Flux-Corrected Remap (FCR) [9] for a collection of classical remap test cases. However, a series of “torture” tests in one and two dimensions demonstrate that FVFT-OBR is significantly more robust than FCR and the representative advection-based remappers. The dual QP provides a convenient structure-exploiting setting for the effective solution of FVFT-OBR by the reflective Newton method [10]. Numerical studies in [1] indicate that the computational cost of FVFT-OBR is proportional, up to a constant, to the cost of an explicit remapper such as FCR. The proportionality constant observed in [1] varies between 1.8 and 3.2.

These figures do not count potential gains from the increased robustness of FVFT-OBR, which enables larger displacements between the old and new mesh. Nonetheless, further efficiency gains in OBR are desirable to enhance its standing as a viable competitor to explicit remappers. Accordingly, the main focus of this paper is on improving the computational efficiency of OBR. Our principal goal is to develop an OBR formulation that fully retains the robustness and accuracy of FVFT-OBR, yet has the computational efficiency of the explicit, state-of-the-art FCR.

The main efficiency bottleneck in FVFT-OBR is the coupled system of linear inequality constraints. The coupling of the variables stems from writing the new cell masses in flux form, which automatically conserves the total mass. While the flux form is needed for conventional remappers, it is arguably less instrumental in an optimization setting where conservation of mass can be treated as an explicit constraint. To take advantage of this fact, we consider an alternative mass-variable mass-target OBR (MVMT-OBR) formulation in which the optimization variables are the net mass updates per cell. While this formulation introduces a single global equality constraint to conserve mass, it completely decouples the inequality constraints because there is only one variable per cell. As a result, by switching to these new variables we change the structure of the OBR problem from a global linear-inequality constrained QP to a singly linearly constrained QP with simple bounds.

Of course, trading automatic mass conservation for an explicit constraint only makes sense if the resulting QP can be solved more efficiently. Using the structure of the MVMT-OBR problem, and the fact that in remap the old and new grids are close, we are able to develop a simple, efficient and easily parallelizable algorithm for the primal MVMT-OBR QP. Preliminary studies of the computational cost suggest that this
algorithm is as efficient as the explicit FCR. This makes the new formulation fully competitive in terms of speed with any conventional remapper. At the same time, thorough computational studies confirm that the MVMT-OBR formulation retains the accuracy and robustness of FVFT-OBR.

The use of the fully decoupled net mass update variables instead of the coupled flux variables opens up an additional possibility to simplify and improve MVMT-OBR by discarding the variables associated with static cells, i.e., the cells that do not move during the rezone step. Since the net mass update on a static cell equals zero, we can increase the efficiency of MVMT-OBR by not computing the variables whose values are known to be zero. This modification could bring about significant additional performance gains in applications such as propagation of waves, in which there are large numbers of static cells. We show that skipping the static cells in the MVMT-OBR formulation does not lead to a loss of key theoretical properties such as existence of optimal solutions and preservation of linearity.

The remainder of this section introduces the relevant notation. Section 2 presents the MVMT-OBR formulation. There we also prove that MVMT-OBR is well-posed and preserves linear functions. The section closes with a discussion of the swept region implementation of MVMT-OBR. Section 3 develops the optimization algorithm for the solution of the MVMT-OBR QP. Numerical studies in Section 4 focus on the accuracy, robustness and efficiency of MVMT-OBR. We summarize our conclusions and map directions for future work in Section 5.

1.1. Notation

The computational domain \( \Omega \) is a bounded subset of \( \mathbb{R}^d \), \( d = 1, 2, 3 \). The old (Lagrangian) grid \( K_h(\Omega) \) is a conforming partition of \( \Omega \) into cells \( \kappa_i, i = 1, \ldots, K \). The total numbers of vertices, edges and sides in the mesh are \( V \), \( E \) and \( S \), respectively. The sets of all vertices, sides and cells in an entity \( \Xi \) are \( V(\Xi) \), \( S(\Xi) \), and \( C(\Xi) \), respectively. For instance, \( S(K_h) \) are the sides in the old mesh and \( V(K_i) \) are the vertices of cell \( \kappa_i \).

When two cells \( \kappa_i \) and \( \kappa_j \) have a common side we label it by \( s_{ij} \). Because \( s_{ij} \) and \( s_{ji} \) represent the same entity in the mesh, to avoid ambiguity we adopt the convention \( i < j \). For instance, referring to the cartoon in Figure 1, we label the side between \( \kappa_5 \) and \( \kappa_2 \) by \( s_{25} \), the side between \( \kappa_5 \) and \( \kappa_6 \) by \( s_{56} \) and so on. This rule induces the orientation of \( s_{ij} \) by selecting the unit normal on the side that points towards the cell with the larger number. Returning to the example in Figure 1, we orient \( s_{25} \) using the unit normal that points into \( \kappa_5 \), while for \( s_{56} \) we use the unit normal that points towards \( \kappa_6 \). The side-to-cell incidence matrix \( D \) is a \( K \times S \) matrix with entries \( d_{i,kl} \) such that \( d_{i,kl} = 0 \) if \( s_{kl} \) is not a side of \( \kappa_i \), \( d_{i,kl} = 1 \) if the unit normal to the side \( s_{kl} \) matches the direction of the outer unit normal to the boundary of \( \kappa_i \), and \( d_{i,kl} = -1 \) otherwise. Succinctly,

\[
 d_{i,kl} = \begin{cases} 
 0 & \text{if } s_{kl} \notin S(\kappa_i) \\
 1 & \text{if } s_{kl} \in S(\kappa_i) \text{ and } i = k \\
 -1 & \text{if } s_{kl} \in S(\kappa_i) \text{ and } i = l 
\end{cases} \tag{1.1} 
\]

![Figure 1: Illustration of the naming and the orientation conventions for cell sides in two dimensions. The sides are \( s_{ij} \) with \( i < j \). The unit normal \( n_{ij} \) on \( s_{ij} \) points away from cell \( \kappa_i \). The unit tangent \( t_{ij} \) is such that the pair \( (t_{ij}, n_{ij}) \) is positively oriented. The blue circles are the signs of the non-zero entries in the row of the side-to-cell incidence matrix corresponding to \( \kappa_5 \).](image)
The blue ovals in Figure 1 illustrate this rule.

In two dimensions we also need to select a unit tangent on each side. We follow the rule that the unit tangent $t_{ij}$ on $s_{ij}$ is such that the pair $(t_{ij}, n_{ij})$ has positive orientation, i.e., $\det([t_{ij} \ n_{ij}]) = 1$.

The new (rezoned) grid $\tilde{K}_h(\Omega)$ is another conforming partition of $\Omega$ into cells $\tilde{\kappa}_i$, $i = 1, \ldots, \tilde{K}$. In this paper we restrict attention to pairs of new and old grids having the same connectivity. Therefore, $K_h(\Omega)$ and $\tilde{K}_h(\Omega)$ have the same numbers of vertices, sides and cells, i.e., $\bar{V} = V$, $\bar{S} = S$ and $\bar{K} = \tilde{K}$. Without loss of generality we assume that the vertices $\bar{v}_i$, the sides $\tilde{s}_{kl}$ and the cells $\tilde{\kappa}_j$ in $\tilde{K}_h(\Omega)$ are numbered by the same numbers as in $K_h(\Omega)$, i.e., their Lagrangian prototypes are $v_i$, $s_{kl}$ and $\kappa_j$, respectively.

The quantities and the entities on the new grid will have the tilde accent, e.g. $\tilde{F}$, whereas the quantities and the entities on $K_h(\Omega)$ will have no accent. For instance,

$$
\mu_i = \int_{\kappa_i} dV \quad \text{and} \quad \tilde{b}_j = \frac{\int_{\tilde{s}_{ij}} x dV}{\mu_j},
$$

are the unsigned measure of the old cell $\kappa_i$ and the barycenter of the new cell $\tilde{\kappa}_j$, respectively. The signed measure\(^3\) of a set $\gamma$ is $\mu^*(\gamma)$.

![Figure 2: Lagrangian cell $\kappa_i$ and its neighborhoods.](image)

The neighborhood $N(\kappa_i)$ of $\kappa_i$ includes the cell $\kappa_i$ itself and all cells that share a vertex (in 1D), vertex or side (in 2D) and vertex, edge or side (in 3D) with $\kappa_i$. The strict neighborhood of $\kappa_i$ is defined by $N'(\kappa_i) = N(\kappa_i) \setminus \kappa_i$. The side neighborhood $N_S(\kappa_i)$ comprises of all cells in $N(\kappa_i)$ which share a side with $\kappa_i$. The subset of cells in $N(\kappa_i)$ that share an edge but not a side with with $\kappa_i$ is the edge neighborhood $N_E(\kappa_i)$ and the subset of cells that shares a vertex but not an edge or a side with $\kappa_i$ is the vertex neighborhood $N_V(\kappa_i)$. The strict versions $N'_S(\kappa_i)$, $N'_E(\kappa_i)$ and $N'_V(\kappa_i)$ of these neighborhoods do not include $\kappa_i$. Figure 2 gives an example of $N(\kappa_i)$, $N'(\kappa_i)$, $N_S(\kappa_i)$ and $N_V(\kappa_i)$ in two dimensions.

We assume that $K_h(\Omega)$ satisfies the locality condition

$$
\tilde{\kappa}_i \subset N(\kappa_i), \quad \text{for all } i = 1, \ldots, \tilde{K},
$$

which makes precise the assumption that $K_h(\Omega)$ and $\tilde{K}_h(\Omega)$ are “close”.

Suppose that $\mathcal{F}$ and $\tilde{\mathcal{F}}$ are some collections of old and new cells, respectively. The notation $i \in \mathcal{F}$, $i \in \tilde{\mathcal{F}}$ means that the index $i$ runs over the indices of the cells in $\mathcal{F}$, resp. $\tilde{\mathcal{F}}$. For instance, $j \in N(\kappa_i)$ means that $j$ loops over the indices of the cells in the neighborhood of $\kappa_i$.

Remark 1.1. Because a side is shared by at most two cells, dimension of $N_S(\kappa_i)$ for interior cells is exactly the number of sides in $\kappa_i$ plus one. In contrast, more than two cells can share an edge in 3D and a vertex in 2D and 3D. As a result, the dimensions of $N_E(\kappa_i)$ (in 3D) and $N_V(\kappa_i)$ (in 2D and 3D) can be arbitrary. Figure 2 shows an example where the number of cells from $N_V(\kappa_i)$ attached to each of the nodes of $\kappa_i$ ranges from 0 to 2.

\(^3\)Reduction of the volume integral to a surface integral and taking into account the surface orientation defines the signed measure of a domain.
2. Formulation of the optimization-based remap through aggregate mass transfer

We begin with a formal statement of the mass-density remap problem. Then we review the flux-variable
flux-target OBR (FVFT-OBR) and proceed to define the mass-variable mass-target OBR (MVMT-OBR).
We show that MVMT-OBR is well-posed and preserves linear densities. Discussion of implementation using
swept region approximation completes the section.

2.1. Statement of the mass-density remap problem

The statement of the mass-density remap problem [11, 9] requires additional notation. The primitive
variable (density) is a scalar function \( \rho(x) > 0 \) on \( \Omega \). The conserved variable is the total mass
\( M = \int_{\Omega} \rho(x) dV = \sum_{i=1}^{K} m_i = \sum_{i=1}^{K} \rho_i \mu_i \)
where

\[
\rho_i = \frac{\int_{\kappa_i} \rho(x) dV}{\mu_i} = \frac{m_i}{\mu_i} \quad \text{and} \quad m_i = \int_{\kappa_i} \rho(x) dV = \rho_i \mu_i, \quad i = 1, 2, \ldots, K, \tag{2.1}
\]
are the mean densities and cell masses on the old mesh, respectively. For every old cell \( \kappa_i \) define

\[
\rho_{\text{min}}^{i} = \begin{cases} \min_{j \in N(\kappa_i)} \{ \rho_j \} & \text{if } \kappa_i \cap \partial \Omega = \emptyset \smallskip \vspace{2pt} \\
\min \left\{ \min_{j \in N(\kappa_i)} \{ \rho_j \}, \min_{x \in N(\kappa_i) \cap \partial \Omega} \rho(x) \right\} & \text{if } \kappa_i \cap \partial \Omega \neq \emptyset \end{cases}, \tag{2.2}
\]

\[
\rho_{\text{max}}^{i} = \begin{cases} \max_{j \in N(\kappa_i)} \{ \rho_j \} & \text{if } \kappa_i \cap \partial \Omega = \emptyset \\
\max \left\{ \max_{j \in N(\kappa_i)} \{ \rho_j \}, \max_{x \in N(\kappa_i) \cap \partial \Omega} \rho(x) \right\} & \text{if } \kappa_i \cap \partial \Omega \neq \emptyset \end{cases}. \tag{2.3}
\]

It is straightforward to check that the mean density trivially satisfies the local bounds

\[
\rho_{\text{min}}^{\text{min}} \leq \rho_i \leq \rho_{\text{max}}^{\text{max}}, \quad i = 1, \ldots, K. \tag{2.4}
\]

The inequalities (2.4) together with (2.1) imply that cell masses satisfy similar local bounds:

\[
m_{\text{min}}^{\text{min}} := \rho_{\text{min}}^{\text{min}} \mu_i \leq m_i \leq \rho_{\text{max}}^{\text{max}} \mu_i =: m_{\text{max}}^{\text{max}}, \quad i = 1, \ldots, K. \tag{2.5}
\]

In the mass-density remap problem we assume that we know the mean density values \( \rho_i \) on the old grid
cells \( \kappa_i \), and the boundary values of \( \rho(x) \) on \( \partial \Omega \). Given this information, we seek accurate approximations \( \tilde{m}_i \) for the masses of the new cells \( \tilde{\kappa}_i \),

\[
\tilde{m}_i \approx \tilde{m}_i^{\text{ex}} = \int_{\tilde{\kappa}_i} \rho(x) dV, \quad i = 1, \ldots, K, \tag{2.6}
\]
such that the following conditions hold:

**C1.** The total mass is conserved:

\[
\sum_{i=1}^{K} \tilde{m}_i = \sum_{i=1}^{K} m_i = M.
\]

**C2.** If \( \rho(x) \) is a global linear function on \( \Omega \), then the remapped masses are exact:

\[
\tilde{m}_i = \tilde{m}_i^{\text{ex}} = \int_{\tilde{\kappa}_i} \rho(x) dV, \quad i = 1, \ldots, K. \tag{2.7}
\]
C3. The approximation of the mean density on the new cells

\[ \tilde{\rho}_i = \frac{\tilde{m}_i}{\tilde{\mu}_i} \]  

(2.8)

satisfies the local bounds

\[ \rho_i^{\text{min}} \leq \tilde{\rho}_i \leq \rho_i^{\text{max}} ; \quad i = 1, \ldots, K, \]  

(2.9)

where (2.2)–(2.3) define \( \rho_i^{\text{min}} \) and \( \rho_i^{\text{max}} \). Equivalently, the following local bounds hold:

\[ \tilde{m}_i^{\text{min}} := \rho_i^{\text{min}} \tilde{\mu}_i \leq \tilde{m}_i \leq \rho_i^{\text{max}} \tilde{\mu}_i =: \tilde{m}_i^{\text{max}}. \]  

(2.10)

\[ \Box \]

2.2. Flux-variable flux-target optimization-based remap

Recall the strict neighborhood

\[ N'(\kappa_i) = N(\kappa_i) \setminus \kappa_i. \]  

The flux-form formula [11]

\[ \tilde{m}_i^{\text{ex}} = m_i + \sum_{j \in N'(\kappa_i)} F_{ij}^{\text{ex}}, \]  

(2.11)

where

\[ F_{ij}^{\text{ex}} = \int_{\tilde{\kappa}_i \cap \kappa_j} \rho(x) dV - \int_{\kappa_i \cap \tilde{\kappa}_j} \rho(x) dV \quad j \in N'(\kappa_i),, \]  

(2.12)

are the exact mass fluxes, holds on any two grids that satisfy (1.3). The substitution of the exact fluxes \( \{F_{ij}^{\text{ex}}\} \) in (2.11) with antisymmetric approximations \( \{F_{ij}\} \), \( F_{ij} = -F_{ji} \) yields the mass-conserving flux-form formula

\[ \tilde{m}_i = m_i + \sum_{j \in N'(\kappa_i)} F_{ij} \]  

(2.13)

for the approximation of the new masses. The discrete fluxes \( \{F_{ij}\} \) are the variables in the FVFT-OBR formulation and the constraints are (i) the antisymmetry \( F_{ij} = -F_{ji} \) of the discrete fluxes and (ii) the local mass bounds (2.10). We enforce antisymmetry directly by using only fluxes \( F_{ij} \) with \( i < j \) and writing the flux sum in (2.13) as

\[ \sum_{j \in N'(\kappa_i)} F_{ij} = \sum_{j \in N'(\kappa_i) \atop i < j} F_{ij} - \sum_{j \in N'(\kappa_i) \atop i > j} F_{ji}. \]

The FVFT-OBR solution minimizes the Euclidean distance between \( F_{ij} \) and some suitable target fluxes \( F_{ij}^T \), \( i < j \), subject to (2.10). Concisely, the FVFT-OBR formulation is the following global inequality-constrained QP [1]:

\[
\begin{aligned}
\text{min}_{F_{ij}} \quad & \frac{1}{K} \sum_{i=1}^{K} \sum_{j \in N'(\kappa_i) \atop i \leq j} (F_{ij} - F_{ij}^T)^2 \\
\text{subject to} \quad & \delta \tilde{m}_i^{\min} \leq \sum_{j \in N'(\kappa_i) \atop i < j} F_{ij} - \sum_{j \in N'(\kappa_i) \atop i > j} F_{ji} \leq \delta \tilde{m}_i^{\max}, \quad i = 1, \ldots, K,
\end{aligned}
\]

(2.14)

where

\[ \delta \tilde{m}_i^{\min} = \tilde{m}_i^{\min} - m_i \quad \text{and} \quad \delta \tilde{m}_i^{\max} = \tilde{m}_i^{\max} - m_i. \]  

(2.15)

A density reconstruction \( \rho^h(x) \) on \( K_h(\Omega) \) that is exact for linear functions defines the target fluxes in (2.14):

\[ F_{ij}^T := \int_{\tilde{\kappa}_i \cap \kappa_j} \rho^h_j(x) dV - \int_{\kappa_i \cap \tilde{\kappa}_j} \rho^h_i(x) dV \quad i < j. \]  

(2.16)
Because our goal is to preserve linear densities, in this paper we restrict attention to piecewise linear density reconstructions, i.e., functions \( \rho^h(x) \) whose restriction \( \rho_i^h(x) = \rho^h|_{\kappa_i} \) to cell \( \kappa_i \) is linear for all \( i = 1, \ldots, K \).

By construction, any optimal solution of QP (2.14) satisfies the local mass bounds (2.10) and conserves the total mass. Because (2.10) and (2.9) are equivalent, the FVFT-OBR solution fulfills \( C_1 \) and \( C_3 \) in Section 2.1. In [1] we prove that FVFT-OBR preserves linear densities by showing that if \( \rho(x) \) is linear, then \( \{ F^T_{ij} \} \) are feasible and \( F_{ij} = F^T_{ij} \) is optimal solution of (2.14). As a result, whenever \( \rho(x) \) is linear,

\[
\tilde{m}_i = m_i + \sum_{j \in N'(\kappa_i)} F^T_{ij} = m_i + \sum_{j \in N'(\kappa_i)} F^{ex}_{ij} = m^{ex}_i, \quad i = 1, \ldots, K.
\]  

(2.17)

Therefore, FVFT-OBR satisfies requirement \( C_2 \) in §2.1.

2.3. Mass-variable mass-target optimization-based remap

Using the flux form (2.13) with antisymmetric discrete fluxes automatically conserves the mass in QP (2.14). Yet, the flux form yields a system of globally coupled inequality constraints. This coupling complicates the solution and creates a performance bottleneck. While the flux form is imperative for conventional remappers, its importance diminishes in the optimization setting, where conservation becomes one of the many possible physically motivated constraints. We take advantage of this fact to replace (2.14) with a QP whose structure is more amenable to fast, parallelizable solution.

In a nutshell, we trade the automatic conservation of mass in (2.13) for a simpler set of box constraints plus a single linear constraint to enforce the conservation of mass explicitly. To this end, instead of (2.13) we start with the mass-form representation of the new cell masses

\[
\tilde{m}^{ex}_i = m_i + \delta m^{ex}_i,
\]

(2.18)

where

\[
\delta m^{ex}_i = \int_{\tilde{\kappa}_i} \rho(x) dV - \int_{\kappa_i} \rho(x) dV
\]

(2.19)

is the mass update between \( \kappa_i \) and \( \tilde{\kappa}_i \). The substitution of \( \delta m^{ex}_i \) in (2.19) by an approximation \( \delta m_i \) yields the mass-form formula for the approximate masses on the new cells:

\[
\tilde{m}_i = m_i + \delta m_i.
\]

(2.20)

The approximate mass updates \( \delta m_i \) are the variables in the mass-target mass-variable optimization-based remap (MVMT-OBR). The constraints on these variables are the linear equality constraint

\[
\sum_{i=1}^{K} \delta m_i = 0,
\]

(2.21)

which is necessary and sufficient for (2.20) to conserve the total mass and the box constraints

\[
\delta m^{\text{min}}_i \leq \delta m_i \leq \delta m^{\text{max}}_i, \quad i = 1, \ldots, K,
\]

(2.22)

which follow from inserting (2.20) into (2.10) and using (2.15).

Remark 2.1. The equality constraint (2.21) holds for the exact mass updates:

\[
\sum_{i=1}^{K} \delta m_i^{ex} = \sum_{i=1}^{K} \int_{\tilde{\kappa}_i} \rho(x) dV - \sum_{i=1}^{K} \int_{\kappa_i} \rho(x) dV = \int_{\Omega} \rho(x) dV - \int_{\Omega} \rho(x) dV = 0.
\]

(2.23)

Using (2.19) in conjunction with a piecewise linear density reconstruction \( \rho^h(x) \) on the old grid defines the mass update targets for the MVMT-OBR formulation:

\[
\delta m^T_i = \int_{\tilde{\kappa}_i} \rho^h(x) dV - \int_{\kappa_i} \rho^h(x) dV.
\]

(2.24)
Suppose that (2.16) defines $F^T_{ij}$. It is easy to see that

$$
\sum_{j \in N'(\kappa_i)} F^T_{ij} = \sum_{j \in N'(\kappa_i)} \left( \int_{\tilde{\kappa}_i \cap \kappa_j} \rho^b_j(x) dV - \int_{\kappa_i \cap \tilde{\kappa}_j} \rho^b_i(x) dV \right) = \left( \sum_{j \in N'(\kappa_i)} \int_{\tilde{\kappa}_i \cap \kappa_j} \rho^b_j(x) dV + \int_{\tilde{\kappa}_i \cap \kappa_j} \rho^b_i(x) dV \right) - \left( \sum_{j \in N'(\kappa_i)} \int_{\kappa_i \cap \tilde{\kappa}_j} \rho^b_i(x) dV + \int_{\kappa_i \cap \tilde{\kappa}_j} \rho^b_i(x) dV \right) = \int_{\tilde{\kappa}_i} \rho^b(x) dV - \int_{\kappa_i} \rho^b(x) dV = \delta m^T_i.
$$

In other words, the target mass update on cell $\kappa_i$ is the sum of the target mass fluxes corresponding to the cells in $N'(\kappa_i)$:

$$
\delta m^T_i = \sum_{j \in N'(\kappa_i)} F^T_{ij}. \quad (2.25)
$$

To sum it up, the MVMT-OBR solution minimizes the Euclidean distance between $\{\delta m_i\}$ and the targets $\{\delta m^T_i\}$, subject to the linear constraint (2.21) and the box constraints (2.22):

$$
\begin{aligned}
& \min_{\delta m_i} \sum_{i=1}^K (\delta m_i - \delta m^T_i)^2 & \text{subject to} \\
& \sum_{i=1}^K \delta m_i = 0 & \text{and} & \delta \tilde{m}^\text{min}_i \leq \delta m_i \leq \delta \tilde{m}^\text{max}_i, \quad i = 1, \ldots, K.
\end{aligned} \quad (2.26)
$$

Problem (2.26) is a singly linearly constrained QP with simple bounds. In contrast to (2.14) the new formulation requires an additional linear equality constraint to conserve the mass, but completely decouples the inequality constraints. In Section 3 we take advantage of this structure and the locality assumption (1.3) to develop fast, efficient and parallelizable optimization algorithm for (2.26).

2.4. Properties of MVMT-OBR

We first establish that (2.26) is a well-posed optimization problem. Then we prove that MVMT-OBR preserves globally linear densities.

**Theorem 2.1.** Assume that the locality condition (1.3) holds for $K_\kappa(\Omega)$ and $\tilde{K}_\kappa(\Omega)$ and that $\{\rho_i\}$ are nonnegative real numbers that specify the mean density on the Lagrangian mesh. Then the QP (2.26) has a unique optimal solution.

**Proof.** Because (2.26) has a strictly convex objective function, it suffices to show that (2.26) has a non-empty feasible set, i.e., that there is at least one set of values $\delta m = (\delta m_1, \ldots, \delta m_K)$, which satisfy the linear constraint (2.21) and the box constraints (2.22). Let

$$
\delta m_i = \sum_{j \in N'(\kappa_i)} \left( \rho_j \mu(\tilde{\kappa}_i \cap \kappa_j) - \rho_i \mu(\kappa_i \cap \tilde{\kappa}_j) \right), \quad i = 1, \ldots, K. \quad (2.27)
$$

We prove that $\delta m$ is feasible. To show (2.21) we break the equality constraint into two parts:

$$
\sum_{i=1}^K \delta m_i = \sum_{i=1}^K \sum_{j \in N'(\kappa_i)} \rho_j \mu(\tilde{\kappa}_i \cap \kappa_j) - \sum_{i=1}^K \sum_{j \in N'(\kappa_i)} \rho_i \mu(\kappa_i \cap \tilde{\kappa}_j) = A - B.
$$

After changing the order of summation in the first part and collecting the like terms we get

$$
A = \sum_{j=1}^K \rho_j \left( \sum_{i \in N'(\kappa_j)} \mu(\tilde{\kappa}_i \cap \kappa_j) \right) = \sum_{j=1}^K \rho_j \left( \sum_{i \in N'(\kappa_j)} \mu(\tilde{\kappa}_i \cap \kappa_j) \pm \mu(\kappa_j \cap \kappa_i) \right) = \sum_{j=1}^K \rho_j \left( \mu(\kappa_j) - \mu(\kappa_j \cap \kappa_i) \right),
$$

$$
B = \sum_{i=1}^K \sum_{j \in N'(\kappa_i)} \rho_i \mu(\kappa_i \cap \tilde{\kappa}_j) = \sum_{i=1}^K \sum_{j \in N'(\kappa_i)} \rho_i \mu(\kappa_i \cap \kappa_j) = \sum_{i=1}^K \rho_i \mu(\kappa_i) = \sum_{i=1}^K \rho_i \mu(\kappa_i),
$$

where
while collecting the like terms in the second part yields
\[
B = \sum_{i=1}^{K} \rho_i \left( \sum_{j \in N(k_i)} \mu(k_i \cap \tilde{k}_j) \right) = \sum_{i=1}^{K} \rho_i \left( \sum_{j \in N(k_i)} \mu(k_i \cap \tilde{k}_j) \pm \mu(k_i \cap k_j) \right) = \sum_{i=1}^{K} \rho_i \left( \mu(k_i) - \mu(k_i \cap k_j) \right).
\]

Therefore, \( A = B \), which proves that \( \delta m \) satisfies the linear equality constraint. To show that \( \delta m \) also satisfies the box constraints, note that (2.27) and \( m_i = \mu \rho(k_i) \) imply
\[
m_i + \delta m_i = \rho_i \left( \mu(k_i) - \sum_{j \in N(k_i)} \mu(k_i \cap \tilde{k}_j) \right) + \sum_{j \in N(k_i)} \rho_j \mu(\tilde{k}_i \cap k_j).
\]
The term in the parenthesis equals \( \mu(k_i \cap k_j) \) and so,
\[
m_i + \delta m_i = \rho_i \mu(k_i \cap \tilde{k}_i) + \sum_{j \in N(k_i)} \rho_j \mu(\tilde{k}_i \cap k_j) = \sum_{j \in N(k_i)} \rho_j \mu(\tilde{k}_i \cap k_j).
\]

On the other hand, (2.2)–(2.3) guarantee that \( \rho_i^{\min} \leq \rho \leq \rho_i^{\max} \) for all cells in \( N(k_i) \). As a result,
\[
m_i + \delta m_i = \sum_{j \in N(k_i)} \rho_j \mu(\tilde{k}_i \cap k_j) \leq \rho_i^{\max} \sum_{j \in N(k_i)} \mu(\tilde{k}_i \cap k_j) = \rho_i^{\max} \mu_i;
\]
\[
m_i + \delta m_i = \sum_{j \in N(k_i)} \rho_j \mu(\tilde{k}_i \cap k_j) \leq \rho_i^{\min} \sum_{j \in N(k_i)} \mu(\tilde{k}_i \cap k_j) = \rho_i^{\min} \mu_i.
\]

Because the box constraints (2.22) are equivalent to
\[
\rho_i^{\min} \mu_i \leq m_i + \delta m_i \leq \rho_i^{\max} \mu_i, \quad i = 1, \ldots, K,
\]
this proves the theorem. \(\square\)

Suppose that the exact density is a globally linear function. The following theorem specializes a result from [1] and establishes sufficient conditions for the QP (2.26) to preserve linear densities.

**Theorem 2.2.** Let \( \rho(x) = c_0 + c^T x \). Assume that \( K_h(\Omega) \) and \( \tilde{K}_h(\Omega) \) satisfy the locality condition (1.3), the density reconstruction \( \rho^h(x) \) is exact for linear functions, and (2.24) defines the target mass updates \( \{\delta m_i^T\} \). Let \( B_i \) denote the set of barycenters of the Lagrangian cells in \( N(k_i) \),
\[
B_i = \{b_j \mid j \in N(k_i)\},
\]
and let \( \tilde{b}_i \) be the barycenter of the rezoned cell \( \tilde{k}_i \). The conditions
\[
\tilde{b}_i \in \mathcal{H}(B_i) \quad \text{if } k_i \cap \partial \Omega = \emptyset, \tag{2.28}
\]
\[
\tilde{b}_i \in \mathcal{H}(B_i \cup (N(k_i) \cap \partial \Omega)) \quad \text{if } k_i \cap \partial \Omega \neq \emptyset, \tag{2.29}
\]
where \( \mathcal{H}(\cdot) \) denotes the convex hull, are sufficient for the remapped masses to be exact, i.e., for \( \tilde{m}_i = \tilde{m}_i^{ex} \) on every rezoned cell \( \tilde{k}_i \).

**Proof.** By assumption \( \rho^h(x) \) is exact for linear functions and so, on every Lagrangian cell \( \rho^h(x) = \rho(x) = c_0 + c^T x \). As a result, \( \delta m_i^T = \delta m_i^{ex} \) and
\[
m_i + \delta m_i^T = \tilde{m}_i^{ex}.
\]

Therefore, to prove \( \tilde{m}_i = \tilde{m}_i^{ex} \) it suffices to show that the targets \( \{\delta m_i^{ex}\} \) are feasible for QP (2.26).

Equation (2.23) in Remark 2.1 confirms that the exact mass updates satisfy the equality constraint (2.21). On the other hand, the bounds (2.22) for \( \{\delta m_i^{ex}\} \) are equivalent to the bounds on the cell masses
\[
\rho_i^{\min} \mu_i \leq \tilde{m}_i^{ex} \leq \rho_i^{\max} \mu_i.
\]

Assuming that (2.28)–(2.29) hold, these bounds follows from [1, Theorem 2.2]. \(\square\)
2.5. Active cell option

We call a new cell \( \tilde{k}_i \) static if \( \tilde{k}_i = k_i \). A cell that is not static is active. In the context of ALE methods static cells arise when the rezoning algorithm, which aims to reduce the grid distortion accrued during the Lagrangian phase, skips cells that it deems to be of a sufficiently high “quality”. There are many practically important applications in which large numbers of cells in \( \Omega \) remain static. A shock or a wave propagating through a domain is one such example. As a rule, the cells outside of a small region where the solution features reside remain static throughout the rezone phase.

Because the mass and the mean density do not change on static cells, that is, \( \tilde{\rho}_i = \rho_i \) and \( \tilde{m}_i = m_i \), it follows that the net mass update per static cell is zero, i.e., \( \delta m_i = 0 \) whenever \( \tilde{k}_i = k_i \). On the other hand, (2.26) is a global optimization problem whose solution may not yield zero mass updates on the static cells. We can improve both the efficiency of MVMT-OBR and the physical quality of its solution by skipping the mass update computations for all static cells. Such a modification is consistent with the notion of remap as a constrained interpolation between two grids: on static cells the field representation does not change and so there is “nothing to interpolate”.

Let \( K_a \) denote the number of active cells in \( K_a(\Omega) \). Without loss of generality assume that the active cells are the first \( K_a \) cells in the mesh. The mass-variable mass-target formulation with an active cell option, MVMT(a)-OBR, restricts the optimization in (2.26) to the net mass updates on the active cells only. Specifically, we set the net mass updates to zero on all static cells, that is,

\[
\delta m_{i} = 0, \quad i = K_a + 1, \ldots, K,
\]

and solve a QP for the net updates on the active cells only:

\[
\begin{align*}
\min_{\delta m_i} & \quad \sum_{i=1}^{K_a} (\delta m_i - \delta m_i^T)^2 \\
\text{subject to } & \quad \sum_{i=1}^{K_a} \delta m_i = 0 \quad \text{and} \quad \delta \tilde{m}_i^\text{min} \leq \delta m_i \leq \delta \tilde{m}_i^\text{max}, \quad i = 1, \ldots, K_a.
\end{align*}
\]

The active cell option does not change the key properties of MVMT-OBR, i.e., the conclusions of Theorem 2.1 and Theorem 2.2 remain in full force for MVMT(a)-OBR.

Let us first confirm that Theorem 2.1 continues to hold for (2.30). Removing the terms that correspond to the static cells from the objective function in (2.26) does not change its strict convexity. Consequently, as before, it suffices to show that MVMT(a)-OBR has a non-empty feasible set. In particular, we will show that (2.27) defines \( \delta m \) that is also feasible for the MVMT(a)-OBR. Suppose that \( k_i \) is a static cell. Since \( \tilde{k}_i = k_i \) it follows that

\[
\mu(\tilde{k}_i \cap k_j) = \mu(k_i \cap k_j) = 0 \quad \text{and} \quad \mu(k_i \cap \tilde{k}_j) = \mu(\tilde{k}_i \cap k_j) = 0.
\]

As a result, formula (2.27) yields

\[
\delta m_i = \sum_{j \in N^s(k_i)} \left( \rho_j \mu(\tilde{k}_i \cap k_j) - \rho_i \mu(\tilde{k}_i \cap k_j) \right) = 0
\]

for all \( \delta m_i \) corresponding to static cells and so, \( \delta m = (\delta m_1, \ldots, \delta m_{K_a}, 0, \ldots, 0) \). Theorem 2.1 implies that \( \delta \tilde{m}_i^\text{min} \leq \delta m_i \leq \delta \tilde{m}_i^\text{max} \) for \( i = 1, \ldots, K_a \), i.e., \( \delta m \) is feasible for MVMT(a)-OBR.

To show that MVMT(a)-OBR preserves linear densities, note that for static cells

\[
\sum_{j \in N^s(k_i)} F^T_{ij} = \sum_{j \in N^s(k_i)} \left( \int_{\tilde{k}_i \cap k_j} \rho_j^h(x) dV - \int_{k_i \cap k_j} \rho_i^h(x) dV \right) = \sum_{j \in N^s(k_i)} \left( \int_{k_i \cap k_j} \rho_j^h(x) dV - \int_{\tilde{k}_i \cap k_j} \rho_i^h(x) dV \right) = 0,
\]

for all \( i = 1, \ldots, K_a \).
and so, (2.25) implies that $\delta m^T = 0$ whenever $\tilde{\kappa}_i = \kappa_i$. Therefore, the proof of linearity preservation in Theorem 2.2 trivially specializes to MVMT(a)-OBR.

Because MVMT-OBR and MVMT(a)-OBR have identical theoretical properties and differ only in the number of variables, in what follows we write both methods as the QP

$$
\begin{align*}
\min_{\delta m_i} & \sum_{i=1}^{K'} (\delta m_i - \delta m^T_i)^2 \\
\text{subject to} & \sum_{i=1}^{K'} \delta m_i = 0 \quad \text{and} \quad \delta \tilde{m}_{i}^{\min} \leq \delta m_i \leq \delta \tilde{m}_{i}^{\max}, \quad i = 1, \ldots, K',
\end{align*}
$$

where $K' = K$ for MVMT-OBR, $K' = K_a$ – the number of active cells, for MVMT(a)-OBR. The last $K - K'$ variables are set to zero: $\delta m_i = 0, \quad i = K' + 1, \ldots, K$.

### 2.6. Swept region approximation

Recall that $\rho^h(x)$ is a piecewise linear function defined with respect to the old mesh. As a result,

$$
\delta m_i^T = \int_{\kappa_i} \rho^h(x) dV - \int_{\kappa_i} \rho^h(x) dV = \sum_{j \in N'(\kappa_i)} \int_{\tilde{\kappa}_i \cap \kappa_j} \rho^h_j(x) dV - \int_{\kappa_i} \rho^h_i(x) dV ,
$$

where $\rho^h_i(x)$ is the restriction of $\rho^h(x)$ to the Lagrangian cell $\kappa_i$. An implementation of formula (2.32) requires the computation of the intersections between the new and old cells. While software tools for this task exist, the swept region approach [11, §4] offers a simpler and more efficient alternative, especially in three dimensions. Swept regions are defined by the movement of the sides of the old cells into a new configuration. As a result, swept regions are completely determined by the coordinates of the old and new cell vertices.

For simplicity, we explain the swept region approach in two dimensions. The sketch in Figure 3 uses the cell numbering from Figure 1 and shows the swept regions resulting from connecting the side vertices of $\tilde{\kappa}_5$ and its Lagrangian prototype $\kappa_5$. Each swept region $\sigma_{ij}$ corresponds to a side $s_{ij}$ in the old mesh. The unit tangent $t_{ij}$ to that side induces orientation on $\sigma_{ij}$. Because swept regions are associated with the sides of

![Figure 3: Swept region nomenclature in two dimensions. Cell and side numbers follow the example in Figure 1. The unit face tangent $t_{ij}$ indicates the orientation of the swept region $\sigma_{ij}$. The blue ovals indicate the signs of the entries in the side-to-cell incidence matrix (1.1).](image-url)

the cells, their use with the flux form (2.13) of the new masses is natural. Specifically, to develop the swept region approximation of FVFT-OBR we restrict the summation in (2.13) to the side neighborhood of $\kappa_i$:

$$
\bar{m}_i = m_i + \sum_{j \in N^c_5(\kappa_i)} F_{ij}
$$

(2.33)
The optimization variables $F_{ij}$ are mass fluxes associated with the swept regions. To define the targets for these variables we consider integration of a piecewise linear density reconstruction $\rho^b$ on $\sigma_{ij}$. As before, $\rho_i^b$ is the linear restriction of $\rho^b$ to the old cell $\kappa_i$.

Note that a swept region $\sigma_{ij}$ can intersect more than one cell. Consequently, exact integration of $\rho^b$ on $\sigma_{ij}$ would also require intersections between the swept region and the old cells. To avoid the computation of these intersections we approximate $\int_{\sigma_{ij}} \rho^b dV$ using either $\rho_i^b$ or $\rho_j^b$ to compute the target flux for $F_{ij}$. The choice of the two possible restrictions of $\rho^b$ depends on (i) the signed area $\mu^*(\sigma_{ij})$ of the swept region, and (ii) the orientation of $s_{ij}$ encoded in the side-to-cell incidence matrix $D$. Specifically, instead of (2.16) we define the target fluxes according to the following formula:

$$F_{ij}^{TS} = \begin{cases} \int_{\sigma_{ij}} \rho_i^b(x) dV & \text{if } \mu^*(\sigma_{ij}) d_{i,ij} < 0 \\ \int_{\sigma_{ij}} \rho_j^b(x) dV & \text{if } \mu^*(\sigma_{ij}) d_{i,ij} > 0 \end{cases}.$$  

(2.34)

Figure 4 illustrates the choice of the reconstruction component as a function of the signed area and the side orientation. The swept region approximation (2.34) is exact for linear functions [12]. As a result, all theoretical conclusions about FVFT-OBR continue to hold when the approximate target fluxes (2.34) are used in lieu of the “true” targets (2.16); see [1].

We now proceed to motivate the swept region approximation for the MVMT-OBR formulation. According to (2.25), when using exact cell intersections the target mass update $\delta m_i^T$ on $\kappa_i$ equals the sum of the target mass fluxes for that cell. This relationship prompts the approximation

$$\delta m_i^{TS} = \sum_{j \in N_5(\kappa_i)} F_{ij}^{TS},$$  

(2.35)

where (2.34) defines the target fluxes $F_{ij}^{TS}$.

It is easy to see that the swept region approximation does not change the theoretical properties of the MVMT-OBR formulation. Theorem 2.1 does not depend on the choice of the targets and so it remains in full force. In contrast, Theorem 2.2 does depend on the target selection. However, because the swept region approximation is exact for linear densities, it follows that formula (2.35) is also exact for such densities, i.e., there holds $\delta m_i^{TS} = \delta m_i^{CS}$, whenever $\rho$ is linear. This is sufficient to carry out the proof of the theorem in the case when (2.35) defines the targets.

### 2.7. Flux-variable mass-target formulation

In this section we briefly mention an alternative QP formulation of remap, which combines the features of FVFT-OBR and MVMT-OBR. We explain why this formulation is less attractive for the development of an optimization-based remap algorithm.
The FVFT-OBR objective minimizes the Euclidean distance between the flux variables and the target fluxes. We can define another optimization objective by combining the flux variables that contribute to the same cell. The resulting QP minimizes the Euclidean distance between the sums of the mass fluxes exchanged between new and old cells and the sums of the corresponding target fluxes, subject to the same bounds on the flux variables as in (2.14):

$$
\min_{F_{ij}} \left( \sum_{i=1}^{K'} \left( \sum_{j \in N'(\kappa_i)} F_{ij} - \sum_{j \in N'(\kappa_i)} F_{Tij} \right) \right)^2 \quad \text{subject to}
$$

$$\delta \tilde{m}_{i}^{\text{min}} \leq \sum_{j \in N'(\kappa_i)} F_{ij} - \sum_{j \in N'(\kappa_i)} F_{ji} \leq \delta \tilde{m}_{i}^{\text{max}} \quad i = 1, \ldots, K'.
$$

(2.36)

From (2.25) we know that the sum of the target fluxes in the objective function of (2.36) gives the mass target $\delta m_{i}^{T}$ of the MVMT-OBR formulation. Furthermore, the sum of the flux variables is effectively a mass update

$$
\delta m_{i} := \sum_{j \in N'(\kappa_i)} F_{ij}
$$

for cell $\kappa_i$. As a result, we can write (2.36) in the following flux-variable mass-target (FVMT) form:

$$
\min_{F_{ij}} \left( \sum_{i=1}^{K'} (\delta m_{i} - \delta m_{i}^{T}) \right)^2 \quad \text{subject to}
$$

$$\delta \tilde{m}_{i}^{\text{min}} \leq \sum_{j \in N'(\kappa_i)} F_{ij} - \sum_{j \in N'(\kappa_i)} F_{ji} \leq \delta \tilde{m}_{i}^{\text{max}} \quad i = 1, \ldots, K'.
$$

(2.37)

The FVMT-OBR formulation combines the flux variables of FVFT-OBR with the objective function of MVMT-OBR. Compared to MVMT-OBR this formulation suffers from two key drawbacks. First, since (2.36) uses flux variables, it inherits the coupled system of linear inequality constraints from FVFT-OBR, which is the main efficiency bottleneck of the FVFT-OBR algorithm. Second, one can show that the objective function of (2.36) has a large null space. Therefore, the FVMT-OBR formulation is not strictly convex and the optimal solution of (2.36) is not unique. This complicates the efficient numerical solution of FVMT-OBR. For these reasons we do not pursue FVMT-OBR in this paper.

3. Optimization Algorithm

In this section we discuss the solution of the optimization problem (2.31), which we restate for ease of reference:

$$
\min_{\delta m_{i}} \sum_{i=1}^{K'} (\delta m_{i} - \delta m_{i}^{T})^2 \quad \text{subject to}
$$

$$\sum_{i=1}^{K'} \delta m_{i} = 0 \quad \text{and} \quad \delta \tilde{m}_{i}^{\text{min}} \leq \delta m_{i} \leq \delta \tilde{m}_{i}^{\text{max}}, \quad i = 1, \ldots, K'.
$$

We remind that $K' = K$ – the number of all cells, in the case of MVMT-OBR, $K' = K_a$ – the number of active cells, in the case of MVMT(a)-OBR and the last $K - K'$ variables are set to zero:

$$
\delta m_{i} = 0; \quad i = K' + 1, \ldots, K.
$$

Optimization problems of this type are known in the optimization literature as the singly linearly constrained quadratic programs with simple bounds. A recent publication with new techniques for their solution is [13], where the authors propose an algorithm that runs in expected linear time and is directly applicable.
to (2.31). While the algorithm proposed here is motivated by [13], it is considerably simpler due to several features of (2.31) that are unique to the context of remap.

A key observation in developing an algorithm for (2.31) is that the related optimization problem without the mass conservation constraint, $\sum_{i=1}^{K'} \delta m_i = 0$, is fully separable. This problem can be solved by independently solving $K'$ one-dimensional quadratic programs with simple bounds, i.e., its solution cost is $O(K')$.

Thus, our goal is to satisfy the remaining constraint $\sum_{i=1}^{K'} \delta m_i = 0$ in a few iterations of cost $O(K')$ each.

We solve (2.31) by a direct application of the Karush-Kuhn-Tucker (KKT) theory, see, e.g., [14, Ch.12]. We define the Lagrangian functional $L : \mathbb{R}^{K} \times \mathbb{R} \times \mathbb{R}^{K'} \times \mathbb{R}^{K'} \rightarrow \mathbb{R}$, as is the vector of primal optimization variables, and $\lambda \in \mathbb{R}$, $\mu_1 \in \mathbb{R}^{K'}$, with $(\mu_1)_i = \mu_{1,i}$, and $\mu_2 \in \mathbb{R}^{K'}$, with $(\mu_2)_i = \mu_{2,i}$, are the Lagrange multipliers. The gradient of the Lagrangian with respect to the primal variables is given by

$$
\frac{\partial}{\partial \delta m_i} L(\delta m, \lambda, \mu_1, \mu_2) = \delta m_i - \delta m_i^T - \lambda \mu_1,i + \mu_2,i, \quad \text{for } i = 1, \ldots, K'.
$$

As shown in Section 2, $\delta m_{i, \min}$ and $\delta m_{i, \max}$ are such that the constraints in (2.31) are consistent, i.e., the optimization problem has a solution. Furthermore, strict convexity yields a unique global minimizer. The necessary and sufficient optimality conditions for (2.31) are:

$$
\delta m_i = \delta m_i^T + \lambda + \mu_{1,i} - \mu_{2,i} \quad i = 1, \ldots, K' \tag{3.1a}
$$

$$
\tilde{\delta m}_{i, \min} \leq \delta m_i \leq \tilde{\delta m}_{i, \max} \quad i = 1, \ldots, K' \tag{3.1b}
$$

$$
\mu_{1,i} \geq 0, \quad \mu_{2,i} \geq 0 \quad i = 1, \ldots, K' \tag{3.1c}
$$

$$
\mu_{1,i} (\delta m_i - \tilde{\delta m}_{i, \min}) = 0, \quad \mu_{2,i} (-\delta m_i + \tilde{\delta m}_{i, \max}) = 0 \quad i = 1, \ldots, K' \tag{3.1d}
$$

$$
\sum_{i=1}^{K'} \delta m_i = 0. \tag{3.1e}
$$

We solve the conditions (3.1) directly. First, we focus on the conditions (3.1a)-(3.1d), which are separable in the index $i$. For any fixed value of $\lambda$ a solution to (3.1a)-(3.1d) is given by

$$
\begin{cases}
\delta m_i = \delta m_i^T + \lambda; & \mu_{1,i} = \mu_{2,i} = 0 \\
\delta m_i = \tilde{\delta m}_{i, \min}; & \mu_{2,i} = 0, \mu_{1,i} = \delta m_i - \delta m_i^T - \lambda \\
\delta m_i = \tilde{\delta m}_{i, \max}; & \mu_{1,i} = 0, \mu_{2,i} = \delta m_i - \delta m_i + \lambda
\end{cases}
$$

for all $i = 1, \ldots, K'$. Ignoring the Lagrange multipliers $\mu_1$ and $\mu_2$ and treating $\delta m_i$ as a function of $\lambda$ yields

$$
\delta m_i(\lambda) = \text{median}(\tilde{\delta m}_{i, \min}, \delta m_i^T + \lambda, \tilde{\delta m}_{i, \max}), \quad i = 1, \ldots, K',
$$

where we note that this is an $O(K')$ computation.

Second, we adjust $\lambda$ in an outer iteration in order to satisfy the constraint $\sum_{i=1}^{K'} \delta m_i(\lambda) = 0$. When we find the $\lambda^*$ such that $\sum_{i=1}^{K'} \delta m_i(\lambda^*) = 0$ holds, we will have solved the optimality conditions (3.1). From (3.3) it is clear that $\sum_{i=1}^{K'} \delta m_i(\lambda)$ is a piecewise linear, monotonically increasing function of a single scalar variable $\lambda$. Therefore, a secant method can be efficiently employed as the outer iteration.
A basic secant approach, such as the one described in [15, p.28-31], typically requires modifications to ensure fast global convergence. The algorithm outlined in [13] tailors such modifications to the context of secant methods applied to general singly linearly constrained QPs with simple bounds. However, in the context of MVMT-OBR it turns out that globalization is unnecessary. This is primarily a consequence of the strict convexity of (2.31) and the quality of the initial guess \( \lambda_0 = 0 \). Substituting \( \lambda_0 = 0 \) into (3.3) yields \( \delta m_i(\lambda_0) = \text{median}(\tilde{\delta}m^\min_i, \tilde{\delta}m^T_i + \lambda_0, \tilde{\delta}m^\max_i) \), for \( i = 1, \ldots, K' \), in other words the solution of the optimization problem (2.31) without the constraint \( \sum_{i=1}^{K'} \delta m_i = 0 \). In Section 4 we demonstrate on several examples that \( \delta m_i(\lambda_0) \), \( i = 1, \ldots, K' \), violate the mass conservation constraint very slightly. This can be traced back to the locality assumption (1.3). In summary, at each remap step we have \( |\sum_{i=1}^{K'} \delta m_i(\lambda_0)| \approx 0 \), so only a few (1-5) iterations of the basic secant method are needed to achieve full feasibility. We also note that since \( \sum_{i=1}^{K'} \delta m_i(\lambda) \) is a piecewise linear function, the secant method always gives a solution that is accurate to machine precision for our choice of stopping tolerances. The complete algorithm is stated below.

Algorithm 1 [Secant method for solving the MVMT-OBR problem, (2.31)]

1. Initialization: Set \( \lambda_0 \leftarrow 0 \), \( \eta \leftarrow 10^{-12} \) and \( \Delta \lambda^{FD} \leftarrow 10^{-8} \).
2. Finite difference step:
   (a) Compute \( \delta m_i(\lambda_0) \leftarrow \text{median}(\tilde{\delta}m^\min_i, \tilde{\delta}m^T_i + \lambda_0, \tilde{\delta}m^\max_i) \) for \( i = 1, \ldots, K' \).
   (b) Compute residual \( r_p \leftarrow \sum_{i=1}^{K'} \delta m_i(\lambda_0) \).
      If \( |r_p| < \eta \), then return \( \delta m_i(\lambda_0) \) for \( i = 1, \ldots, K' \) and stop.
   (c) Compute residual \( r_c \leftarrow \sum_{i=1}^{K'} \delta m_i(\lambda_0 + \Delta \lambda^{FD}) \).
3. While \( |r_c| > \eta \) (Secant Iteration)
   (a) Compute \( \delta m_i(\lambda_c) \leftarrow \text{median}(\tilde{\delta}m^\min_i, \tilde{\delta}m^T_i + \lambda_c, \tilde{\delta}m^\max_i) \) for \( i = 1, \ldots, K' \).
   (b) Compute residual \( r_c \leftarrow \sum_{i=1}^{K'} \delta m_i(\lambda_c) \).
   (c) Set \( \alpha \leftarrow (\lambda_p - \lambda_c)/(r_p - r_c) \). Set \( r_p \leftarrow r_c \).
   (d) Set \( \lambda_p \leftarrow \lambda_0 \). Set \( \lambda_c \leftarrow \lambda_p - \alpha r_p \).
4. Return \( \delta m_i(\lambda_c) \) and stop.

4. Computational studies

4.1. Preservation of shape, monotonicity and linearity

In this section we present numerical examples that demonstrate that FVFT-OBR and MVMT-OBR give virtually identical results when subjected to the “torture” tests developed in [1]. The overall purpose of these tests is to numerically confirm the theoretical properties of the methods in extreme scenarios. For completeness we include the results of the same studies performed with a flux-corrected remap (FCR) algorithm. For all algorithms we use swept regions for the target flux reconstruction.

The first test examines the ability of the algorithms to preserve the shape of a given density function. The test involves a one-dimensional mesh consisting of three cells. In the original or “old” configuration the cells are of equal lengths. In the remapped or “new” configuration the middle cell is compressed by a factor of approximately 6.6, see [1] for the detailed setup. The density function has the shape of an asymmetrical peak. An accurate method should preserve the shape of the peak on the new mesh. It is evident from
Figure 5 that FVFT-OBR and MVMT-OBR preserve the peak, giving identical results, while FCR loses accuracy due to the local approximation of the global FVFT-OBR formulation.

![Original Density (Old Mesh)](image1)

![FCR Result (New Mesh)](image2)

![FVFT-OBR Result (New Mesh)](image3)

![MVMT-OBR Result (New Mesh)](image4)

Figure 5: FVFT-OBR and MVMT-OBR preserve the shape of the peak, giving identical results, while FCR loses accuracy and transforms the peak into a step function.

The second test focuses on the preservation of monotonicity for a given density function. We use a simple two-dimensional extension of the previous one-dimensional experiment. The same setup is used in [1]. As in one dimension, the original mesh is uniform, while in the new mesh the middle cell is compressed equally in both spatial directions, see Figure 6. The density function is linear. We report loss of monotonicity if the bounds in (2.14) or (2.26) are violated after a single remap step. We note that the compressive mesh motion illustrated in Figure 6 is permitted according to the locality assumption (1.3) and the linearity preservation conditions (2.28)-(2.29) established for FVFT-OBR and MVMT-OBR. Table 1 confirms that FVFT-OBR and MVMT-OBR preserve monotonicity, which is a direct consequence of the construction of the schemes as constrained optimization algorithms. At the same time, as explained in [1], the condition on mesh motion for the monotonicity of the swept-region donor-cell method [11, p.279] is violated by this example. This defect is inherited by FCR, resulting in the loss of monotonicity for high compressions of the middle cell.

![Figure 6: A 3×3 uniform mesh (left pane) and a “compressed” mesh (right pane) with a 4×4-fold compression of the middle cell.](image5)

The third test examines whether the methods preserve linear density functions. We use the same torture mesh as in the second experiment. The \( L_1 \) error is evaluated after two remap steps (original mesh → compressed mesh → original mesh). Since the linearity preservation conditions (2.28)-(2.29) are satisfied for the compressive mesh motion, we expect that FVFT-OBR and MVMT-OBR preserve linear densities regardless of how much the middle cell is compressed. This is confirmed in Table 2, where FVFT-OBR and MVMT-OBR give very similar results. In contrast, FCR fails to preserve linear densities at the compression
Table 1: Monotonicity of FVFT-OBR, MVMT-OBR and FCR, implemented using swept regions, with respect to the remap of a linear density function in two dimensions, for different compression ratios $\ell \times \ell : 1$ of the middle cell. FVFT-OBR and MVMT-OBR are monotone throughout, while FCR is not.

<table>
<thead>
<tr>
<th>$\ell$</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>14</th>
<th>15</th>
<th>16</th>
<th>100</th>
</tr>
</thead>
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<tr>
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<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>MVMT-OBR</td>
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<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>FCR</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
<td>no</td>
</tr>
</tbody>
</table>

ratios of $4 \times 4 : 1$ and above.

Table 2: $L_1$ errors in the FVFT-OBR, MVMT-OBR and FCR remap of a linear density function in two dimensions, for different compression ratios $\ell \times \ell : 1$ of the middle cell. Errors close to machine precision are highlighted. FVFT-OBR and MVMT-OBR preserve linear densities for arbitrarily compressed middle cells, while FCR does not.

<table>
<thead>
<tr>
<th>$\ell$</th>
<th>3</th>
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<th>5</th>
<th>15</th>
<th>16</th>
<th>100</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVFT-OBR</td>
<td>(L$_1$ err) 1.02e-16 3.02e-16 2.22e-16 6.17e-16 2.22e-15 1.26e-13</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MVMT-OBR</td>
<td>(L$_1$ err) 6.78e-17 3.05e-16 2.25e-16 1.54e-15 1.70e-15 1.46e-13</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FCR</td>
<td>(L$_1$ err) 9.25e-17 3.01e-03 7.68e-03 3.28e-02 3.49e-02 1.51e-01</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4.2. Asymptotic accuracy

In this section we compare the asymptotic accuracy of FVFT-OBR and MVMT-OBR in the context of a continuous rezone strategy. The appropriate notions of remap error and convergence rates can be defined through cyclic remap tests [11]. The methodology is described in great detail in [1, 12, 11].

We consider two cyclic mesh motions from [1]: the smooth tensor-product motion, resulting in a sequence of slowly evolving rectangular meshes, and the repeated-repair motion, giving a sequence of rapidly changing rectangular meshes, derived from the torture motion illustrated in Figure 6. For the convergence studies involving the tensor-product motion, we use the sine, peak and shock densities referenced in [1, 11]. We test convergence under the repeated-repair motion using the sine density.

For the repeated-repair motion, the absolute accuracy (and convergence rate) of MVMT-OBR equals that of FVFT-OBR, see Table 4. For the tensor-product motion, FVFT-OBR has a slight advantage in terms of absolute accuracy, see Table 3. As shown in Section 4.6 this accuracy can be easily matched by MVMT-OBR when using slightly finer computational meshes.

4.3. Robustness

The purpose of the test in this section is to subject the FVFT-OBR and MVMT-OBR algorithms to mesh motions that may violate the locality assumption (1.3) and the linearity preservation conditions (2.28)-(2.29). In particular, we examine the behavior of the methods on $64 \times 64$ tensor-product cyclic meshes as the number of remaps, $R$, is decreased much beyond the previously chosen value of $R = 320$. Equivalently, we may say that the so-called pseudo-time step, $1/R$, is increased significantly beyond $1/R = 1/320$. For completeness, we include the FCR results from [1].

It is evident from Table 5 that FVFT-OBR and MVMT-OBR perform similarly, in other words they are both robust when subjected to mesh deformations that are beyond the theoretical guarantees of the methods. In contrast, the FCR algorithm violates linearity preservation early, and eventually becomes unstable.
<table>
<thead>
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<th>L₁ rate</th>
<th>L₁ err</th>
<th>L₁ rate</th>
<th>L₁ err</th>
<th>L₁ rate</th>
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</thead>
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<td>4.91e-04</td>
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<td>2.55e-03</td>
<td>—</td>
<td>2.88e-02</td>
<td>—</td>
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<td>6.16e-05</td>
<td>3.00</td>
<td>8.90e-04</td>
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<td>3.10e-04</td>
<td>1.52</td>
<td>1.06e-02</td>
<td>0.72</td>
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<tr>
<td>512×512</td>
<td>2560</td>
<td>9.89e-07</td>
<td>2.98</td>
<td>1.09e-04</td>
<td>1.52</td>
<td>6.35e-03</td>
<td>0.73</td>
</tr>
</tbody>
</table>

**Table 3:** FVFT-OBR and MVMT-OBR L₁ errors and convergence rate estimates for the sine, peak and shock densities using 4 tensor-product cyclic meshes. The convergence rates are very similar. The absolute errors are slightly lower for FVFT-OBR.

<table>
<thead>
<tr>
<th>#cells</th>
<th>#remaps</th>
<th>L₁ err</th>
<th>L₁ rate</th>
<th>L₁ err</th>
<th>L₁ rate</th>
<th>L₁ err</th>
<th>L₁ rate</th>
</tr>
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<td>—</td>
<td>2.88e-03</td>
<td>—</td>
<td>3.03e-02</td>
<td>—</td>
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<tr>
<td>128×128</td>
<td>640</td>
<td>8.09e-05</td>
<td>2.98</td>
<td>1.08e-03</td>
<td>1.42</td>
<td>1.83e-02</td>
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<tr>
<td>256×256</td>
<td>1280</td>
<td>1.04e-05</td>
<td>2.97</td>
<td>4.04e-04</td>
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<td>1.10e-02</td>
<td>0.73</td>
</tr>
<tr>
<td>512×512</td>
<td>2560</td>
<td>1.32e-06</td>
<td>2.97</td>
<td>1.51e-04</td>
<td>1.42</td>
<td>6.65e-03</td>
<td>0.73</td>
</tr>
</tbody>
</table>

**Table 4:** FVFT-OBR and MVMT-OBR L₁ errors and convergence rate estimates for the sine density using 4 repeated-repair cyclic meshes. The convergence rates are very similar. The absolute errors are slightly lower for FVFT-OBR.

<table>
<thead>
<tr>
<th>R = 213</th>
<th>R = 212</th>
<th>R = 211</th>
<th>R = 155</th>
<th>R = 154</th>
<th>R = 153</th>
<th>R = 100</th>
<th>R = 50</th>
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<tbody>
<tr>
<td>FVFT-OBR</td>
<td>1.32e-13</td>
<td>1.42e-13</td>
<td>1.60e-13</td>
<td>4.60e-09</td>
<td>4.06e-06</td>
<td>1.53e-05</td>
<td>1.97e-03</td>
</tr>
<tr>
<td>MVMT-OBR</td>
<td>1.32e-13</td>
<td>1.42e-13</td>
<td>1.60e-13</td>
<td>4.59e-09</td>
<td>2.07e-06</td>
<td>7.53e-06</td>
<td>1.27e-03</td>
</tr>
<tr>
<td>FCR</td>
<td>1.32e-13</td>
<td>5.32e-08</td>
<td>1.10e-06</td>
<td>2.26e-03</td>
<td>2.35e-03</td>
<td>2.44e-03</td>
<td>5.73e+04</td>
</tr>
</tbody>
</table>

**Table 5:** L₁ errors in the FVFT-OBR, MVMT-OBR and FCR remap of a linear density function on the 64×64 tensor-product cyclic mesh, for different values of the pseudo-time step 1/R. Errors smaller than 1e-8 are highlighted. The OBR methods fail to preserve linear densities at R = 154, while FCR fails at R = 212, resulting in a pseudo-time step advantage for FVFT-OBR and MVMT-OBR of 212/154 ≈ 1.4. Beyond this point, the OBR methods exhibit a graceful loss of accuracy; FCR becomes numerically unstable.

### 4.4. Locality

An interesting question is whether the optimization-based remap algorithms respect the notion of “locality” or “causality”, which is a feature of purely local, albeit less accurate and less robust methods like FCR. To examine the behavior of OBR methods, we construct an example in which only a single mesh vertex is displaced, and measure the mass updates computed by the methods on every cell. We note that if FCR is applied to such single-vertex motion, the mass updates will be zero everywhere except in the immediate neighborhood of the vertex (a four-cell neighborhood in this case).

The single-vertex motion is defined by moving the center vertex of the mesh, with coordinates \((x, y)\), to the position \((x + h \cos(\alpha), y + h \sin(\alpha))\), where \(\alpha = 23\pi/180\) and \(h \in \{1/16, 1/32, 1/64, 1/128\}\), for the mesh sequence \(\{16×16, 32×32, 64×64, 128×128\}\), respectively. The remapped density function is a symmetric Gaussian of amplitude 1 with standard deviation 0.1, centered at \((0.5, 0.5)\). We plot the magnitudes of mass updates, \(\delta m_i\), after a single remap step. The mass updates are scaled by \(K\), the total number of cells.
While MVMT-OBR generates nonzero mass updates everywhere, we observe from Figure 7, top pane, that there is a sharp drop-off in the magnitudes of mesh updates between the four-cell neighborhood surrounding the displaced vertex and the remainder of the domain. The size of the drop-off increases as the mesh is refined. For the 128×128 mesh, see Figure 8, bottom pane, the mass updates are approximately seven orders of magnitude smaller for all cells outside of the four-cell neighborhood of the displaced vertex. Thus we may argue that MVMT-OBR is an “asymptotically” local method.

In contrast, FVFT-OBR generates mass updates that are considerably more localized, see Figure 7, middle pane. Nonetheless, the behavior of FVFT-OBR is not identical to that of FCR, as eight cells, instead of four, contribute with non-zero mass updates. Thus we may label FVFT-OBR “weakly” local.

Finally, we examine the behavior of the mass-variable mass-target algorithm with the active cell option, MVMT(a)-OBR. We recall that MVMT(a)-OBR eliminates the mass-update variables that correspond to static cells. Figure 7 shows that this algorithmic modification yields an optimization-based remap scheme with purely local behavior, when applied to the single-vertex mesh motion. We recommend the use of MVMT(a)-OBR whenever significant portions of the computational domain are static.

4.5. Conservation properties

This section examines the numerical behavior of MVMT-OBR with respect to mass conservation. Our test is designed to verify a claim made in Section 3, namely that the mass updates $\delta m_i(\lambda_0)$, $i = 1, \ldots, K$, where $\lambda_0 = 0$ is the initial Lagrange multiplier guess, violate the mass conservation constraint only slightly. We demonstrate that this is true in a cumulative sense (and so, clearly, at each remap step). We also show that only a handful of iterations of our secant algorithm are needed to fully restore mass conservation.

<table>
<thead>
<tr>
<th>Original mass $\sum_{i=1}^{K} m_i$</th>
<th>% diff</th>
<th>Peak % diff</th>
<th>Shock % diff</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sine</td>
<td>1.0000000000</td>
<td>—</td>
<td>1.0654579749</td>
</tr>
<tr>
<td>Final mass $\sum_{i=1}^{K} m_i(\lambda_0)$</td>
<td>1.0000082955</td>
<td>0.0008</td>
<td>1.0663039593</td>
</tr>
<tr>
<td>Final mass $\sum_{i=1}^{K} m_i(\lambda^*)$</td>
<td>1.0000000000</td>
<td>0</td>
<td>1.0654579749</td>
</tr>
<tr>
<td>Avg. Secant Iterations</td>
<td>1</td>
<td>3</td>
<td>5</td>
</tr>
</tbody>
</table>

Table 6: A comparison of the total mass (i) on the original mesh, (ii) as computed by the “faulty” (nonconservative) MVMT-OBR algorithm after 320 remaps and (iii) as computed by the proper MVMT-OBR algorithm, Algorithm 1, after 320 remaps. The average number of secant iterations for the proper MVMT-OBR algorithm, rounded to the nearest integer, is also reported.

4.6. Computational efficiency

Before proceeding to the comparisons of MVMT-OBR, FVFT-OBR and FCR, it is important to point out that the MVMT-OBR algorithm, Algorithm 1, possesses several advantages over the FVFT-OBR scheme. The FVFT-OBR algorithm relies on a dual QP formulation and a reflective Newton method for its solution, see [1]; the MVMT-OBR algorithm, based on the secant method, is much simpler to implement. The average number of iterations needed by the FVFT-OBR scheme (2-15) is about twice the number of iterations observed in using MVMT-OBR (1-5). Each iteration of FVFT-OBR entails the solution of a sparse but very large, $O(K \times K)$, linear system; each iteration of MVMT-OBR solves $K$ fully decoupled single-variable quadratic programs. Finally, the parallelization of FVFT-OBR necessitates the parallelization of the linear system solve; MVMT-OBR is trivially parallelized and requires minimal communication.
Figure 7: The magnitudes of mass updates scaled by the number of cells in the mesh, $K|\delta m|$, for one step of OBR algorithms subject to the single-vertex motion. The remapped density function is a symmetric Gaussian of amplitude 1 with standard deviation 0.1, centered at (0.5, 0.5). The top pane shows the mass updates generated by MVMT-OBR for the 16×16 and 32×32 meshes, resulting in an “asymptotically” local method. The middle pane shows the mass updates generated by FVFT-OBR – a “weakly” local method; the bottom pane shows the mass updates generated by MVMT(a)-OBR – a local method.
Figure 8: The magnitudes of mass updates scaled by the number of cells in the mesh, $K|\delta m_i|$, for one step of MVMT-OBR subject to the single-vertex motion, shown for the mesh sequence \{16×16, 32×32, 64×64, 128×128\}, across the (0,0)-(1,1) diagonal of the domain.

Table 7 compares the computational cost of FCR, MVMT-OBR and FVFT-OBR for a sequence of cyclic meshes and three target densities.\textsuperscript{4} It is evident that the cost of MVMT-OBR is virtually identical to that of FCR. Additionally, they are on average twice as fast as FVFT-OBR. Table 8 addresses an observation made in Section 4.2, namely that FVFT-OBR may exhibit better absolute accuracy in comparison with MVMT-OBR. Therefore, in order to make the cost comparisons fair, we adjust the mesh resolution and the number of remaps for MVMT-OBR, given in parentheses, to match the $L_1$ error of FVFT-OBR. The computational cost of MVMT-OBR is still significantly lower than the cost of FVFT-OBR.

5. Conclusions

We have developed a new optimization-based conservative, local bounds and linearity-preserving remapping method – MVMT-OBR. Contrary to the existing flux-based remapping methods, which consider mass fluxes exchanged between the new and old mesh cells, MVMT-OBR is based on aggregate mass transfer, that is, the primary unknowns in the optimization formulation are the net mass updates between the new and old cells. The optimization setting lets us treat mass conservation as one of the constraints. The choice of primary variables leads to a singly linearly constrained quadratic optimization problem with simple bounds. The structure of the optimization problem and the fact that in remap the old and new meshes are close allow us to develop an efficient and easily parallelizable optimization algorithm.

We have demonstrated numerically that MVMT-OBR is “asymptotically” local, that is, the domain of influence of a mesh change is limited, as the mesh size goes to zero, to its immediate neighborhood. We have also presented a version of MVMT-OBR with the “active cell option” – MVMT(a)-OBR, which preserves all key features of MVMT-OBR yet treats limited changes in the mesh much like explicit, purely local remappers. We recommend the use of MVMT(a)-OBR whenever a large number of cells are static.

Numerical studies demonstrate that the new algorithm is as accurate and robust as the previously developed flux-based optimization algorithm – FVFT-OBR, but has the same computational cost as the state-of-the-art explicit flux-corrected remapping method – FCR. As future work we plan to develop an algorithm for the recovery of mass fluxes from the aggregate mass updates. This may be needed, for

\textsuperscript{4}All experiments are performed in Matlab\textsuperscript{TM}, Version 7.11.0.584 (R2010b) 64-bit; as in [1], we note that the computational cost of our Matlab\textsuperscript{TM} implementation will be close to that of a Fortran or C implementation, due to a consistent vectorization of algebraic operations.
<table>
<thead>
<tr>
<th># cells</th>
<th># remaps</th>
<th>FCR time (sec)</th>
<th>MVMT-OBR time (sec)</th>
<th>ratio</th>
<th>FVFT-OBR time (sec)</th>
<th>ratio</th>
</tr>
</thead>
<tbody>
<tr>
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<td></td>
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<td></td>
<td></td>
</tr>
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</table>

Table 7: Comparison of the computational costs of FCR, MVMT-OBR and FVFT-OBR, as measured by Matlab™ wall-clock times in seconds, on a single Intel Xeon X5680 3.33GHz processor, for sine, peak and shock densities and the tensor-product cyclic grid. Ratios of run times of MVMT-OBR and FVFT-OBR with respect to FCR are included. The cost of MVMT-OBR is almost identical to the cost of FCR, while FVFT-OBR is on average two times slower.

<table>
<thead>
<tr>
<th># cells</th>
<th># remaps</th>
<th>FVFT-OBR L₁ error</th>
<th>time (sec)</th>
<th>MVMT-OBR L₁ error</th>
<th>time (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>64×64 (71×71)</td>
<td>320 (350)</td>
<td>4.91e-4</td>
<td>6.8</td>
<td>4.94e-4</td>
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<tr>
<td>128×128 (142×142)</td>
<td>640 (705)</td>
<td>6.16e-5</td>
<td>48.7</td>
<td>6.04e-5</td>
<td>33.8</td>
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<tr>
<td>256×256 (283×283)</td>
<td>1280 (1410)</td>
<td>7.82e-6</td>
<td>384.6</td>
<td>7.76e-6</td>
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</tr>
<tr>
<td>512×512 (566×566)</td>
<td>2560 (2825)</td>
<td>9.89e-7</td>
<td>3677.5</td>
<td>9.93e-7</td>
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<tr>
<td>Peak</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>320 (350)</td>
<td>2.55e-3</td>
<td>7.7</td>
<td>2.56e-3</td>
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<td>8.88e-4</td>
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<td>4410.5</td>
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<td>Shock</td>
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<td></td>
</tr>
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<td>64×64 (70×70)</td>
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<td>2.87e-2</td>
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<td>128×128 (138×138)</td>
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<tr>
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<td>3117.1</td>
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<td>2612.2</td>
</tr>
</tbody>
</table>

Table 8: Comparison of the computational costs of FVFT-OBR and MVMT-OBR, as measured by Matlab™ wall-clock times in seconds, on a single Intel Xeon X5680 3.33GHz processor, for sine, peak and shock densities and the tensor-product cyclic grid. In this example the mesh resolution and the number of remaps for MVMT-OBR, given in parentheses, are adjusted to match the L₁ error of FVFT-OBR. The cost of MVMT-OBR is still significantly lower than the cost of FVFT-OBR.

Example, in the context of remapping multiple tracers in climate applications. Another possible extension is a high-order (for example, quadratic-preserving) remapping algorithm, which will require the development of new local bounds that take into account the presence of local extrema.
Acknowledgments

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References