CAM-CM: A Signal Deconvolution Tool for In Vivo Dynamic Contrast-enhanced Imaging of Complex Tissues

Li Chen1,5, Tsung-Han Chan1,2,*, Peter L. Choyke3, Elizabeth M. C. Hillman4, Chong-Yung Chi2, Zaver M. Bhujwalla5, Ge Wang6, Sean S. Wang7, Zsolt Szabo5, Yue Wang1,†
1Bradley Department of Electrical and Computer Engineering and 4School of Biomedical Engineering and Sciences, Virginia Tech, Arlington, VA 22203, USA, 2Department of Electrical Engineering, National Tsing Hua University, Taiwan, ROC, 3Molecular Imaging Program, National Cancer Institute, NIH, Bethesda, MD 20892, USA, 4Department of Biomedical Engineering and Radiology, Columbia University, New York, NY 10027, USA, 5Department of Radiology and Radiological Science, Johns Hopkins University School of Medicine, Baltimore, MD 21205, USA, 7Science, Mathematics, Computer Science House, Poolesville High School, Poolesville, MD 20837, USA.

ABSTRACT
Summary: In vivo dynamic contrast-enhanced imaging tools provide noninvasive methods for analyzing various functional changes associated with disease initiation, progression, and responses to therapy. The quantitative application of these tools has been hindered by its inability to accurately resolve and characterize targeted tissues due to spatially-mixed tissue heterogeneity. CAM-CM (Convex Analysis of Mixtures - Compartment Modeling) signal deconvolution tool has been developed to automatically identify pure-volume pixels located at the corners of the clustered pixel time series scatter simplex and subsequently estimate tissue-specific pharmacokinetic parameters. CAM-CM can dissect complex tissues into regions with differential tracer kinetics at pixel-wise resolution and provide a systems biology tool for defining imaging signatures predictive of phenotypes.

Availability: The MATLAB source code can be downloaded at the authors’ website www.cbil.ece.vt.edu/software.htm
Contact: yuewang@vt.edu

1 INTRODUCTION
In vivo dynamic contrast-enhanced imaging tools provide noninvasive methods for analyzing various functional changes associated with disease initiation, progression, and responses to therapy (McDonald and Choyke, 2003). Typical modalities include dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) (Costouros, et al., 2002), dynamic contrast enhanced optical imaging (Hillman, et al., 2007), positron emission tomography (Zhou, et al., 1997), and spectroscopic computed tomography (Anderson, et al., 2010). These tools exploit the dynamics of contrast accumulation and washout to produce functionally relevant images of vascular perfusion and permeability, metabolism, or gene expression, and can potentially test novel hypotheses and predict drug efficacy.

However, due to spatially-mixed tissue heterogeneity, the precise imaging-based phenotyping by these tools has been hindered by its inability to accurately resolve and characterize targeted functional tissue compartments (Hillman and Moore, 2007). This indistinction between contributions of different tissues to the mixed tracer signals could significantly confound subsequent pharmacokinetics compartmental modeling (CM) (Zhou, et al., 1997) and affect the accuracy of genotype-phenotype association studies (Costouros, et al., 2002; Segal, et al., 2007).

We developed convex analysis of mixtures – compartment modeling (CAM-CM) signal deconvolution tool that enables geometrically- principled, unsupervised, and accurate characterization and delineation of major functional tissue structures from dynamic contrast-enhanced imaging data, not only dissecting complex tissue into regions with differential tracer kinetics at pixel-wise resolution but also substantially improving tissue-specific pharmacokinetic parameter estimation (Wang, et al., 2010). CAM-CM is supported by a well-grounded mathematical framework, and combines the advantages of multivariate clustering, convex geometry analysis, and compartmental modeling. The algorithm possesses a novel, powerful feature allowing pure-volume pixels to be readily identified from the measured pixel time series, without any knowledge of the associated compartmental pharmacokinetics, leading to a completely unsupervised approach. We provide CAM-CM software as an open-source standalone MATLAB application.

2 DESCRIPTION
2.1 Method and Software

Under the framework of compartment modeling in dynamic contrast-enhanced imaging studies, the spatial-temporal signals $x_i(t)$ of tracer concentration at pixel $i$ can be expressed as a non-negative linear combination of the latent tissue-specific compartmental time activity curves $a(t)$, weighted by the relative tissue type proportions $K_j(i)$ at that pixel: $x_i(t) = a_1(t)K_1(i) + \ldots + a_J(t)K_J(i)$, where $J$ is the number of functional tissue compartments (Hillman, et al., 2007). This falls neatly within the definition of a convex set:

$$\mathcal{X} = \left\{ \sum_{j=1}^{J} K_j(i) a_j \mid a_j \in A, K_j(i) \geq 0, \sum_{j=1}^{J} K_j(i) = 1, i = 1, \ldots, N \right\},$$

where $a_i$ is the vector notation of $a(t)$ over time. We have shown that the corner points of the pixel time series convex hull $\mathcal{H}(X)$ correspond to the pure-volume pixels for each tissue compartments (Wang, et al., 2010).

The flowchart of CAM-CM algorithm is given in Fig. 1. The three core components of CAM-CM software include: (1) initialization-free multi-variate clustering of pixel time series into an optimum number of representative and robust clusters using affinity propagation clustering and expectation-maximization mixture model fitting (Frey and Dueck, 2007); (2) convex analysis of mixtures that automatically identifies the pure-volume pixel clusters geometrically located at the corners of the clustered pixel time-series scatter simplex via minimum-error-margin convex-hull-to-data fitting (Wang, et al., 2010); and (3) compartment modeling that estimates tissue-specific pharmacokinetic parameters using only pure-volume pixel time series. The CAM-CM software is implemented in MATLAB, and runs successfully on both Microsoft Windows and Linux platforms.

The CAM-CM software takes input the .mat data files that record the pixel time series of dynamic contrast-enhanced images in matrices. Each row corresponds to a time frame and each column corresponds to a pixel. Running CAM-CM software is automatic and convenient, with only two user-controlled parameters: the number of tissue types $J$ and the sampling time interval between two consecutive dynamic image frames. Results of CAM-CM are provided to the users via a multiplatform graphical summary package, to be a very useful tool for the exploratory analysis of dynamic functional imaging data in many research and clinical applications. We would expect the CAM-CM method, with publicly available open-source software package, to be a very useful tool for the exploratory analysis of dynamic functional imaging data in many research and clinical applications. We plan to develop and deliver future versions of the software written in both R and Java languages.

CAM-CM has been tested on real dynamic contrast-enhanced imaging data. Using DCE-MRI dataset of an advanced breast cancer case (McDonald and Choyke, 2003), CAM-CM analysis reveals two biologically interpretable vascular compartments with distinct kinetic patterns: fast clearance in peripheral “rim” and slow clearance in inner “core” (Fig. 2a-c) which otherwise could not be seen if tissue heterogeneity (84% in this case) was not taken into account, plausibly consistent with the previously reported heterogeneity within tumors (Costouros, et al., 2002). Since angiogenesis is essential to tumor development, it has been widely observed that active angiogenesis in advanced breast tumors often occurs in the peripheral “rim” with co-occurrence of inner-core hypoxia, due to the defective endothelial barrier function and outgrowth blood supply (McDonald and Choyke, 2003).

In another application to dynamic fluorescence molecular imaging data acquired on a mouse after bolus injection of indocyanine green dye (Hillman and Moore, 2007), CAM-CM provides physiologically interpretable biodistribution dynamics of the major organs. Ten fluorescence time-courses (Fig. 2d) show distinct patterns of circulating, accumulating, or metabolizing the dye in different organs, which agree with expected physiological trends, such as late uptake by adipose tissue and fast clearance from the brain region. The merged and color-coded maps of these dissected tissue compartments constitute anatomical structures of the mouse that agree well with a digital anatomical mouse atlas, allowing the longitudinal identification of the internal organs (Fig. 2e) (Hillman and Moore, 2007).

Detailed descriptions on method, simulation-based validation, and more biomedical case studies are included in the supplementary information.

3 DISCUSSION

CAM-CM spatial-temporal analysis can be very effective at revealing multi-compartment structure within dynamic contrast-enhanced imaging data of complex tissues, estimating tissue-specific kinetic parameter values, and characterizing the roles of different functional tissue compartments. We would expect the CAM-CM method, with publicly available open-source software package, to be a very useful tool for the exploratory analysis of dynamic functional imaging data in many research and clinical applications. We plan to develop and deliver future versions of the software written in both R and Java languages.

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REFERENCES


