Recent advances in Computer Vision and Experimental Neuroscience provided insights into mechanisms underlying invariant object recognition. However, due to the different research aims in both fields models tended to evolve independently. A tighter integration between computational and empirical work may contribute to cross-fertilized development of (neurobiologically plausible) computational models and computationally defined empirical theories, which can be incrementally merged into a comprehensive brain model. After reviewing theoretical and empirical work on invariant object perception, this article proposes a novel framework in which neural network activity and measured neuroimaging data are interfaced in a common representational space. This enables direct quantitative comparisons between predicted and observed activity patterns within and across multiple stages of object processing, which may help to clarify how high-order invariant representations are created from low-level features. Given the advent of columnar-level imaging with high-resolution fMRI, it is time to capitalize on this new window into the brain and test which predictions of the various object recognition models are supported by this novel empirical evidence.

Keywords: object perception, view-invariant object recognition, neuroimaging, large-scale neuromodeling, (high-field) fMRI, multimodal data integration

INTRODUCTION

One of the most complex problems the visual system has to solve is recognizing objects across a wide range of encountered variations. Retinal information about one and the same object can dramatically vary when position, viewpoint, lighting, or distance change, or when the object is partly occluded by other objects. In Computer Vision, there are a variety of models using alignment, invariant properties, or part-decomposition methods (Roberts, 1965; Fukushima, 1982; Marr, 1982; Ullman et al., 2001; Viola and Jones, 2001; Lowe, 2004; Torralba et al., 2008), which are able to identify objects across a range of viewing conditions.

Some computational models are clearly biologically inspired and take for example the architecture of the visual system into account (e.g., Wersing and Körner, 2003), or cleverly adapt the concept of a powerful Computer Vision algorithm (e.g., the Fourier-Mellin transform) to a neurobiologically plausible alternative (Sountsov et al., 2011). Such models can successfully detect objects in sets of widely varying natural images (Torralba et al., 2008) and achieve impressive invariance (Sountsov et al., 2011). In general however, computer vision models are developed for practical image analysis applications (handwriting recognition, face detection, etc.) for which fast and accurate object recognition and not neurobiological validity is pivotal. Therefore, these models are generally less powerful in explaining how object constancy arises in the human brain. Indeed, “Models are common; good theories are scarce” as suggested by Stevens (2000, p. 1177). Humans are highly skilled in object recognition, and they outperform machines in object recognition tasks with great ease (Fleuret et al., 2011). This is partly because they are able to strategically use semantics and information from context or memory. In addition, they can direct attention to informative features in the image, while ignoring distracting information. Such higher cognitive processes are difficult to implement, but improve object recognition performance when taken into account (Lowe, 2000). Computer vision models might become more accurate in recognizing objects across a wide range of variations in image input, when implementing algorithms derived from neurobiological observations.

Reciprocally, our interpretation of such neurobiological findings might be greatly improved by insights in the underlying computational mechanisms. Humans can identify objects with great speed and accuracy, even when the object percept is degraded, occluded or presented in a highly cluttered visual scene (e.g., Thorpe et al., 1996). However, which computational mechanisms enable such remarkable performance is not yet fully understood. To create a comprehensive theory of human object recognition and how it achieves invariant object recognition, computational mechanisms derived from modeling efforts should be incorporated in neuroscientific theories based on experimental findings.

In the current paper, we highlight recent developments in object recognition research and put forward a “Common Brain Space” framework (CBS; Goebel and De Weerd, 2009; Peters et al.,
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Exploring Invariant Object Recognition in the Human Visual System

Object recognition, discrimination, and identification are complex tasks. Different encounters with an object are unlikely to take place under identical viewing conditions, requiring the visual system to generalize across changes. Information that is important to retrieve object identity should be effectively processed, while unimportant view-point variations should be ignored. That is, the recognition system should be stable yet sensitive (Marr and Nishihara, 1978), leading to inherent tradeoffs. How the visual system is able to accomplish this task with such apparent ease is not yet understood. There are two classes of theories on object recognition. The first suggests that objects can be recognized by cardinal (“non-accidental”) properties that are relatively invariant to the objects’ appearance (Marr, 1982; Biederman, 1987). Thus, these invariant properties and their spatial relations should provide sufficient information to recognize objects regardless of their viewpoint. However, how such cardinal properties are defined and recognized in an invariant manner is a complex issue (Tarr and Bülthoff, 1995). The second type of theory suggests that there are no such invariants but that objects are stored in the view as originally encountered (which, in natural settings encompasses multiple views being sampled in a short time interval), thereby maintaining view-dependent shape and surface information (Edelman and Bülthoff, 1992). Recognition of an object under different viewing conditions is achieved by either computing quality matches between the input and stored presentations (Perrett et al., 1998; Riesenhuber and Poggio, 1999) or by transforming input to match the view specifications of the stored representation (Bülthoff and Edelman, 1992). The latter normalization can be accomplished by interpolation (Poggio and Edelman, 1990), mental transformation (Tarr and Pinker, 1989), or alignment (Ullman, 1989).

These theories make very different neural predictions. View-invariant theories suggest that the visual system recognizes objects using a limited library of non-accidental properties, and neural representations are invariant. Evidence for such invariant object representations have been found at final stages of the visual pathway (Quiróga et al., 2005; Freiwald and Tsao, 2010). In contrast, the second class of theories assumes that neural object representations are view-dependent, with neurons being sensitive to object transformations. Clearly, the early visual system is sensitive to object appearance: the same object can elicit completely different, non-overlapping neural activation patterns when presented at different locations in the visual field. So, object representations are input specific at initial stages of processing, whereas invariant representations emerge in final stages. However, how objects are represented by intermediate stages of this processing chain is not yet well understood. Likely, multiple different transforms are (perhaps in parallel) performed at these stages. This creates multiple object representations, in line with the various types of information (such as position and orientation) that have to be preserved for interaction with objects. Moreover, position information aids invariant object learning (Einhäuser et al., 2005; Li and DiCarlo, 2008, 2010) and representations can reflect view-dependent and view-invariant information simultaneously (Franzius et al., 2011).

The following section reviews evidence from monkey neurophysiology and human neuroimaging on how object perception and recognition are implemented in the primate brain. As already alluded to above, the visual system is hierarchically organized in more than 25 areas (Felleman and Van Essen, 1991) with initial processing of low-level visual information by neurons in the thalamus, striate cortex (V1) and V2; and of more complex features in V3 and V4 (Carlson et al., 2011). Further processing of object information in the human ventral pathway (Ungerleider and Haxby, 1994), involves higher-order visual areas such as the lateral occipital cortex (LOC; Malach, 1995) and object selective areas for faces (“FFA”; Kanwisher et al., 1997), bodies (“EBA”; Downing et al., 2001), words (“VWFA”; McCandliss et al., 2003), and scenes (“PPA”; Epstein et al., 1999).

The first studies on the neural mechanisms of object recognition were neurophysiological recordings in monkeys. In macaque anterior inferotemporal (IT) cortex, most of the object-selective neurons are tuned to viewpoint-position (Logothetis et al., 1995; Booth and Rolls, 1998), in line with viewpoint-dependent theories. On the other hand, IT neurons also turned out to be more sensitive to changes in “non-accidental” than to equally large pixel-wise changes in other shape features (“metric properties”; Kayaert et al., 2003), providing support for structural description theories (Biederman, 1987). Taken together, these studies provide neural evidence for both theories (see also Rust and DiCarlo, 2010). However, to which degree object representations are stored in an invariant or view-dependent manner across visual areas, and how these representations arise and are matched to incoming information, remains elusive.

Also human neuroimaging studies have not provided conclusive evidence. In fMRI studies, the BOLD signal reflects neural activity at the population rather than single-cell level. The highest functional resolution provided by standard 3 Tesla MRI scanners is around $2 \times 2 \times 2 \text{mm}^3$, which is too coarse to zoom into the functional architecture within visual areas. However, more subtle information-patterns can be extracted using multi-voxel pattern analysis (MVPA; Haynes et al., 2007) or fMRI-adaptation (fMRI-A; Grill-Spector and Malach, 2001). MVPA can reveal subtle differences in distributed fMRI patterns across voxels resulting from small biases in the distributions of differentially tuned neurons that are sampled by each voxel. By using classification techniques developed in machine learning, distributed spatial patterns of different classes (e.g., different objects) can be success-fully discriminated (see Fuentemilla et al., 2010 for a temporal pattern classification example with MEG). For example, changing the position of an object significantly changes patterns in LOC, even more than replacing an object (at the same position) by an object of a different category (Sayres and Grill-Spectro, 2008). Rotating the object (up to 60°) did not change LOC responses however (Eger et al., 2008) suggesting that LOC representations might be view-dependent in only some aspects. fMRI-A exploits the fact that the neuronal (and the corresponding hemodynamic) response is weaker for repeated compared to novel stimuli (Miller and Desimone, 1994). Thus, areas are sensitive to view-dependent...
changes when their BOLD response returns to its initial level for objects that are presented a second time, but now from a different view-point. This technique revealed interesting and unexpected findings. For example, a recent study observed viewpoint and size dependent coding at intermediate processing stages (V4, V3A, MT, and V7), whereas responses in higher visual areas were viewpoint-invariant (Konen and Kastner, 2008). Remarkably, these view-invariant representations were not only found in the ventral (e.g., LOC), but also in the dorsal pathway (e.g., IPS). The dorsal “where/how” or “perception-for-action” pathway is involved in visually guided actions toward objects rather than in identifying objects—which is mainly performed by the ventral or “what” pathway (Goodale and Milner, 1992; Ungerleider and Haxby, 1994). For this role, maintaining viewpoint dependent information in higher dorsal areas seems important, which however was thus not confirmed by the view-invariant results in IPS (but see James et al., 2002). Likewise, another recent study (Dilks et al., 2011) revealed an unexpected tolerance for mirror-reversals in visual scenes in a parahippocampal area thought to play a key role in navigation (e.g., Janzen and van Turenout, 2004) and reorientation (e.g., Epstein and Kanwisher, 1998), functions for which view-dependent information is essential. Furthermore, mixed findings have been reported for the object-selective LOC. For example, different findings on size, position, and viewpoint-invariant representations in different subparts of the LOC have been found (Grill-Spector et al., 1999; James et al., 2002; Vuilleumier et al., 2002; Valyear et al., 2006; Dilks et al., 2011). These divergent findings might be partly related to intricacies inherent to the fMRI-A approach (e.g., Krekelberg et al., 2006), and its sensitivity to the design used (Grill-Spector et al., 2006) and varying attention (Vuilleumier et al., 2005) and task demands (e.g., Ewbank et al., 2011). The latter should not be regarded as obscuring confounds however, since they appear to strongly contribute to our skilled performance. Object perception is accompanied by cognitive processes supporting fast (e.g., extracting the “gist” of a scene, attentional selection of relevant objects) and accurate (e.g., object-verification, semantic interpretation) object identification for subsequent goal-directed use of the object (e.g., grasping; tool-use). These processes engage widespread memory- and frontoparietal attention-related areas interacting with object processing in the visual system (Corbetta and Shulman, 2002; Bar, 2004; Ganis et al., 2007). As the involvement of such top-down processes might be particularly pronounced in humans—and weaker or even absent in monkeys and machines respectively—efforts to integrate computational modeling with human neuroimaging remain essential (see Tagamets and Horwitz, 1998; Corchs and Deco, 2002 for earlier work).

With the advent of ultra-high field fMRI (≥7 Tesla scanners), both the sensitivity (due to increases in signal-to-noise ratio linearly dependent on field strength) and the specificity (due to a stronger contribution of gray-matter microvasculature compared to large draining veins and less partial volume effects) of the acquired signal improves significantly, providing data at a level of detail which previously was only available via invasive optical imaging in non-human species. The functional visual system can be spatially sampled in the range of hundreds of microns, which is sufficient to resolve activation at the cortical column (Yacoub et al., 2008; Zimmermann et al., 2011) and layer (Polimeni et al., 2010) level. Given that cortical columns are thought to provide the organizational structure forming computational units involved in visual feature processing (Hubel and Wiesel, 1962; Tanaka, 1996; Mountcastle, 1997), the achievable resolution at ultra-high fields will therefore not only produce more detailed maps, but really has the potential to yield new vistas on within-area operations.

INTEGRATION OF COMPUTATIONAL AND EXPERIMENTAL FINDINGS IN CBS

The approach we propose is to project the predicted activity in a modeled area onto corresponding cortical regions where empirical data are collected (Figure 1). By interfacing empirical and simulated data in one anatomical “brain space”, direct and quantitative mutual hypothesis testing based on predicted and observed spatiotemporal activation patterns can be achieved. More specifically, modeled units (e.g., cortical columns) are 1-to-1 mapped to corresponding neuroimaging units (e.g., voxels, vertices) in the empirically acquired brain model (e.g., cortical gray matter surface). As a result, a running network simulation creates spatiotemporal data directly on a linked brain model, enabling highly specific and accurate comparisons between neuroimaging and neurocomputational data in the temporal as well as spatial domain. Note that in CBS (as implemented in Neurolator 3D; Goebel, 1993), computational and neuroimaging units can flexibly represent various neural signals (e.g., fMRI, EEG, MEG, fNIRS, or intracranial recordings). Furthermore, both hidden and output layers of the neural network can be projected to the brain model, providing additional flexibility to the framework as predicted and observed activations can be compared at multiple selected processing stages simultaneously (see Figure 2 for an example).

To model the human object recognition system, we developed large-scale networks of cortical column units, which dynamics can either reflect the spike activity, integrated synaptic activity, or oscillating activity (when modeled as burst oscillators), resulting from excitatory and inhibitory synaptic input. To create simulated spatiotemporal patterns, each unit of a network layer (output and/or hidden) is linked to a topographically corresponding patch on a cortical representation via a so-called Network-to-Brain Link (NBL). Via this link, activity of modeling units in the running network is transformed into timecourses of neuroimaging units, spatially organized in an anatomical coordinate system. Importantly, when simulated and measured data co-exist in the same representational space, the same analysis tools (e.g., MVPA, effective connectivity analysis) can be applied to both data sets allowing for quantitative comparisons (Figure 2). See Peters et al. (2010) for further details.

We propose that such a tight integration of neuroimaging and modeling data allows reciprocal fine-tuning and facilitates hypothesis testing at a mechanism level as it leads to falsifiable predictions that can subsequently be empirically tested. Importantly, there is a direct topographical correspondence between computational (cortical columnar) units at the model and brain level. Moreover, comparisons between simulated and empirical data are not limited to activity patterns in output stages
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**FIGURE 1** The Common Brain Space framework: measured neuroimaging data (left panel) and simulated data (right panel) are projected to the same anatomical brain space via network-brain links (see **Figure 2**).

**FIGURE 2** Data Integration in Common Brain Space. Input: (A) Visualization of Common Brain Space (CBS) in Neurolator: Each computational unit of a neural network layer is separately connected to a topographically corresponding location on the cortical sheet via a Network–Brain Link (NBL). In this example, model layers V1, LOC, and FFA are connected to the corresponding brain regions V1, LOC, and FFA on a mesh reconstruction of an individual’s gray-white matter boundary. For this participant, V1, LOC, and FFA were localized using standard retinotopy and related fMRI Region-of-Interest mapping techniques. By connecting a running neural network, activity in the connected layers is projected to the cortical sheet via the NBLs, creating spatially specific timecourses. (B) In Neurolator, functional MRI data can be projected on the cortical mesh, similar to the standard functional-anatomical data co-registration applied in fMRI analyses. Output: (C) Depending on display mode, cortical patches (i.e., vertices) either represent the empirical or the simulated fMRI data. Since the observed and simulated datasets are in the same anatomical space, identical fMRI analyses tools can be used to analyze observed and simulated timeseries.

(i.e., object-selective areas in anterior IT such as FFA or even more anterior in putative “face exemplar” regions; Kriegeskorte et al., 2007), but also at intermediate stages (such as V4 and LOC). Interpreting the role of feature representations at intermediate stages may be essential for a comprehensive brain model of object recognition (Ullman et al., 2002).

Studying several stages of the visual hierarchy simultaneously, by quantitatively comparing ongoing visual processes across
Studies both within and between the simulated and empirically acquired dataset, may help to clarify how higher-order invariant representations are created from lower-level features in several ways. Firstly, this may reveal how object-coding changes along the visual pathway. Incoming percepts might be differently transformed and matched to stored object representations at several stages, with view-dependent matching at intermediate stages and matching of only informative properties (Biederman, 1987; Ullman et al., 2001) at later stages. Secondly, monitoring activity patterns at multiple processing stages simultaneously is desirable, given that early stages are influenced by processing in later stages. To facilitate object recognition, invariant information is for example fed back from higher to early visual areas (Williams et al., 2008), suggesting that object perception results from a dynamic interplay between visual areas. Finally, it is important to realize that such top-down influences are not limited to areas within the classical visual hierarchy, but also engage brain-wide networks involved in “initial guessing” (Bar et al., 2006), object selection (Serences et al., 2004), context integration (Graboi and Lisman, 2003; Bar, 2004), and object verification (Ganis et al., 2007). Such functions should be incorporated in computational brain models to fully comprehend what makes human object recognition so flexible, fast, and accurate. Modeling higher cognitive functions is in general challenging, but may be aided by considering empirical observations in object perception studies where the level of top-down processing varies (e.g., Ganis et al., 2007). The interactions between the visual pathway and frontoparietal system revealed by such fMRI studies can be compared at multiple processing stages to simulations, allowing a more subtle, process-specific fine-tuning of the modeled areas.

A number of recent fMRI studies applied en- and decoding techniques developed in the field of Machine Learning and Computer Vision, to interpret their data (Kriegeskorte et al., 2008; Miyawaki et al., 2008; Haxby et al., 2011; Naselaris et al., 2011; see LaConte, 2011 for an extension to Brain-Computer-Interfaces), showing that both fields are starting to approach each other. For example, by summarizing the complex statistical properties of natural images using a computer vision technique, a visual scene percept could be successfully reconstructed from fMRI activity (Naselaris et al., 2009). The trend to investigate natural vision is noteworthy, given that processing cluttered and dynamic natural visual input rather than artificially created isolated objects poses additional challenges to the visual system (Einhäuser and König, 2010). We believe that now columnar-level imaging is in reach with the advent of high-resolution fMRI (in combination with the recently developed en- and decoding fMRI methods) the time has come to more directly integrate computational and experimental neuroscience, and test which predictions of the various object recognition models are supported by this new type of empirical evidence.

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