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# Learning Laplacians in Chebyshev Graph Convolutional Networks

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## Abstract

*Spectral graph convolutional networks (GCNs) are particular deep models which aim at extending neural networks to arbitrary irregular domains. The principle of these networks consists in projecting graph signals using the eigen-decomposition of their Laplacians, then achieving filtering in the spectral domain prior to back-project the resulting filtered signals onto the input graph domain. However, the success of these operations is highly dependent on the relevance of the used Laplacians which are mostly handcrafted and this makes GCNs clearly sub-optimal. In this paper, we introduce a novel spectral GCN that learns not only the usual convolutional parameters but also the Laplacian operators. The latter are designed "end-to-end" as a part of a recursive Chebyshev decomposition with the particularity of conveying both the differential and the non-differential properties of the learned representations – with increasing order and discrimination power – without overparametrizing the trained GCNs. Extensive experiments, conducted on the challenging task of skeleton-based action recognition, show the generalization ability and the outperformance of our proposed Laplacian design w.r.t. different baselines (built upon handcrafted and other learned Laplacians) as well as the related work.*

designing convolutions on general graph structures [46–49].

The difficulty in extending CNNs to irregular graphs stems from the eclectic properties of graphs and their heterogeneous topological properties which make convolutions *ill-posed*. A particular class of neural machines, known as graph convolutional networks (GCNs), has emerged and seeks to generalize convolutions to irregular graph structures [50]. Their principle consists in learning representations by aggregating signals through nodes and their neighbors, prior to apply convolutions on the resulting aggregates [27, 29, 50, 55, 64–66, 72]. Two categories of GCNs exist in the literature; spatial and spectral. In spatial GCNs [51–56], the representation of a given node is obtained by averaging the representations of its neighbors before applying convolutions using the inner product. In spectral GCNs [15, 28, 46, 47, 49, 57–61], convolutions proceed differently by first projecting filters and input graph signals using the eigen-decomposition of their Laplacians, prior to achieve filtering, and then back-projecting the resulting filtered signals onto the input graph domain [62, 63]. Whereas spectral GCNs make convolutions well defined compared to spatial ones, their drawback resides in the non-localized aspect of the learned filters and also the high complexity of Laplacian eigen-decomposition.

## 1. Introduction

Deep learning is currently achieving a significant progress in computer vision and several related fields [38–41, 67–71]. The purpose of deep learning is to train multi-layered and highly nonlinear parametric models whose inputs correspond to vectorial data (images, etc.) and outputs to their classification or regression [44, 45]. One of the most popular models include convolutional neural networks (CNNs) which operate by shifting equivariant filters and measuring their responses across different image locations. While these operations are well defined on regular grids (namely images), their extension to irregular domains (such as skeletons in action recognition [20–22]) requires

Other spectral GCNs have been introduced in the literature including Chebyshev networks [47] which consider instead localized convolutional filters using a recursive polynomial decomposition. The success of these GCNs is highly reliant on the choice of the Laplacian operators, and most of the existing ones are handcrafted or designed upon the inherent properties of the data. These properties correspond to preexisting (intrinsic) node-to-node relationships such as connectivity in 3D skeletons, links in social networks, protein connectivity in biological systems, etc. Nevertheless, handcrafted Laplacians are not sufficient in order to capture all node relationships as their design is oblivious to the tasks at hand. Indeed, in many real-world applications such as skeleton-based recognition, intrinsic relationships

are suitable to identify individuals using their anthropometric measurements while other extrinsic characteristics are more useful to recognize their dynamics and actions. Hence, extrinsic node-to-node relationships should be inferred in order to maximize the performances of the targeted applications (see Fig. 3). In other words, connectivity in Laplacians should be appropriately learned by including not only the available (intrinsic) node-to-node connections in graphs but also their inferred (extrinsic) relationships.

### 1.1. Related work

Laplacian — or equivalently graph — inference is generally ill-posed, NP-hard [1–3] and most of the existing approaches rely on constraints (including similarity, smoothness, sparsity, band-limitedness, etc.) for a better conditioning [4, 5, 73–86]. Particularly in GCNs, early methods [6, 49, 52] rely on predetermined node-to-node relationships using similarities or the inherent properties of the targeted applications in order to define Laplacian operators. However, in spite of being relatively effective, the potential of these operators is not fully explored as their design is either agnostic to the tasks at hand or achieved using the tedious cross validation. More recent advances aim at defining graph topology — and hence the underlying Laplacian — that best fits a given task [7–12, 15, 59]. For instance, the work in [11] proposes a graph network for semi-supervised classification that learns graph topology, with sparse structure, given a cloud of points; node-to-node connections are modeled with a joint probability distribution on Bernoulli random variables whose parameters are found using bi-level optimization. A computationally more efficient variant is introduced in [12] using a weighted cosine similarity and edge thresholding.

Other solutions make improvement w.r.t. the original GCNs [49] by exploiting symmetric matrices [15] and discovering hidden structural relations (unspecified in the original graphs), using a so-called residual graph adjacency matrix, and by learning a distance function over nodes. The work in [59] introduces a dual architecture with two parallel graph convolutional layers sharing the same parameters. This method considers a normalized adjacency matrix and a positive pointwise mutual information matrix to capture node co-occurrences through random walks sampled from graphs. In the particular context of skeleton-based action recognition, GCNs have been increasingly used [13, 14, 24–26, 35, 36, 42, 43] as they explicitly model, with a better interpretability, the spatial and temporal connectivity among joints. However, while all skeleton-joints contribute in motion, only a few of them are actually relevant to recognize the targeted action categories; hence, other work focuses on learning more complete spatial and temporal joint co-occurrences for skeleton data [15, 30, 34, 37].

### 1.2. Contribution

In this paper, we introduce a novel framework that learns graph topology and Laplacians as a part of GCN and Chebyshev basis design. This basis is expressed using an efficient recursive form evaluated on a single shared Laplacian which conveys both the differential and non-differential properties of the learned graph representations. This Chebyshev basis also captures the statistical properties of the learned representations, with increasing order and discrimination power, without increasing the actual number of training parameters in the resulting GCNs. Different settings of our design are considered including symmetry and orthogonality that further constrain the learned Laplacians and enhance the generalization capacity of our GCNs. Experiments, conducted on the challenging task of skeleton-based human action and hand gesture recognition, show the high accuracy and the outperformance of our method w.r.t. different baselines as well as the related work.

Our proposed method in this paper is different, from all the aforementioned related work, at least in two aspects; on the one hand, none of this related work considers Laplacian learning as a part of Chebyshev basis design. On the other hand, existing methods consider multiple independent matrix operators that capture the actual topology of the input graphs and increase the discrimination power of the learned GCN representations, but this comes at the expense of overparametrized networks and the risk of overfitting. In contrast, our Chebyshev basis design increases the discrimination power of the representations (that capture different hops in graphs), without overparametrizing the trained networks, as the learned Laplacian parameters are shared through all the Chebyshev polynomials. Besides, making the Chebyshev basis<sup>1</sup> orthogonal acts as a regularizer that controls the actual number of training parameters and enhances further the generalization power of our GCNs as corroborated later in experiments.

## 2. Chebyshev Convolutional Networks

Given a graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$  with  $|\mathcal{V}| = n$ ,  $|\mathcal{E}|$  being respectively the number of its vertices and edges, and  $\mathbf{L}$  the Laplacian of  $\mathcal{G}$ . For instance,  $\mathbf{L}$  could be the random walk Laplacian defined as  $\mathbf{L} = \mathbf{I}_n - \mathbf{A}[\mathbf{D}^{-1}(\mathbf{A})]$  where (i)  $\mathbf{I}_n$  is an  $n \times n$  identity matrix, (ii)  $\mathbf{A}$  an adjacency matrix with each entry  $\mathbf{A}_{uu'} > 0$  iff  $(u, u') \in \mathcal{E}$  and 0 otherwise, and (iii)  $\mathbf{D}(\mathbf{A})$  a diagonal degree matrix with each entry  $[\mathbf{D}(\mathbf{A})]_{uu} = \sum_v \mathbf{A}_{vu}$ . Let  $\mathbf{U}\mathbf{\Lambda}\mathbf{U}^\top$  be the eigen-decomposition of  $\mathbf{L}$ , with  $\mathbf{U}$ ,  $\mathbf{\Lambda}$  being respectively the matrix of eigenvectors (graph Fourier basis) and the

<sup>1</sup>The generative aspect of our basis makes it possible to capture different hops of neighbors without increasing the actual number of training parameters and this enhances the discrimination power of the learned representation as shown through this paper.

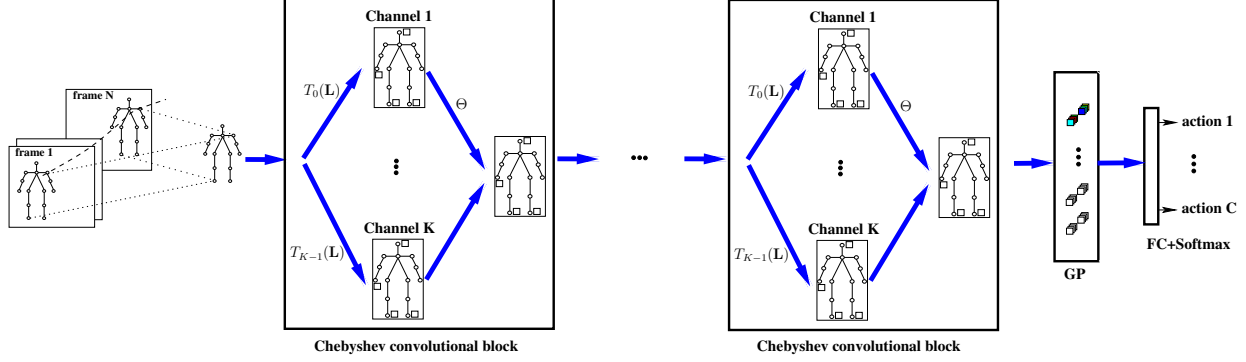


Figure 1. This figure shows the architecture of our Chebyshev Convolutional Network. In each convolutional block, the Chebyshev basis  $\{T_k(\cdot)\}_k$  is first evaluated on the Laplacian  $\mathbf{L}$ , then multiplied by the input graph signal  $\psi(\mathcal{V})$ , and finally aggregated using the parameters  $\theta$ . These Chebyshev convolutional blocks are followed by global average pooling prior to softmax classification. Note that the Laplacian  $\mathbf{L}$  is shared across the basis  $\{T_k(\cdot)\}_k$  and through the Chebyshev convolutional blocks; in practice, only a very few convolutional blocks are necessary as the basis  $\{T_k(\cdot)\}_k$  already captures different hops/depths in graphs. **(Better to zoom the pdf).**

diagonal matrix of its eigenvalues; spectral graph convolution is a well defined operator (see for instance [50]) which is achieved by first projecting a given graph signal  $\psi(\cdot)$  using the eigen-decomposition of  $\mathbf{L}$ , and then multiplying the resulting projection by a convolutional filter prior to back-project the result in the original signal space.

Formally, the convolutional operator  $\star_{\mathcal{G}}$  (rewritten for short as  $\star$ ) on a given graph signal  $\psi(\mathcal{V}) \in \mathbb{R}^{s \times n}$  is  $(\psi \star g_{\theta})_{\mathcal{V}} = \mathbf{U} g_{\theta}(\Lambda) \mathbf{U}^{\top} \psi(\mathcal{V})^{\top}$ ; here  $\top$  is the matrix transpose operator and  $g_{\theta}$  denotes a non-parametric convolutional filter defined as  $g_{\theta}(\Lambda) = \text{diag}(\theta)$  with  $\theta \in \mathbb{R}^n$ . As this filter is not localized, we consider instead [47]

$$(\psi \star g_{\theta})_{\mathcal{V}} = \sum_{k=0}^{K-1} \theta_k T_k(\mathbf{L}) \psi(\mathcal{V})^{\top}, \quad (1)$$

with  $\theta = (\theta_1 \dots \theta_K)^{\top} \in \mathbb{R}^K$  being the learned convolutional filter parameters and  $T_k$  the  $k$ -th order Chebyshev polynomial recursively defined as  $T_k(\mathbf{L}) = 2\mathbf{L} \circ T_{k-1}(\mathbf{L}) - T_{k-2}(\mathbf{L})$ , with  $T_k(\mathbf{L}) \in \mathbb{R}^{n \times n}$ ,  $T_0(\mathbf{L}) = \mathbf{I}_n$ ,  $T_1(\mathbf{L}) = \mathbf{L}$  and  $\circ$  the Hadamard (element-wise) matrix product. When  $\mathbf{L}$  is the combinatorial Laplacian, we consider in practice a rescaled version as  $2\mathbf{L}/\lambda_{\max} - \mathbf{I}_n$  (instead of  $\mathbf{L}$  with  $\lambda_{\max}$  being the largest eigenvalue of  $\mathbf{L}$ ) in order to guarantee the orthogonality of the basis  $\{T_k(\mathbf{L})\}_k$ ; see again [47] and later (in section 3.2) the general rescaling of any Laplacian that guarantees orthogonality of the Chebyshev basis. The whole architecture of this convolution is described in Fig. 1.

### 3. Our Laplacian Design

Considering the tensor of the Chebyshev polynomials  $\{T_k(\mathbf{L})\}_k$  and the loss  $\mathcal{L}$  associated to a given classification task, we turn the design of the Laplacian  $\mathbf{L}$  (thereby the Chebyshev basis  $\{T_k(\mathbf{L})\}_k$ ) as a part of GCN training.

Considering the gradient of  $\mathcal{L}$  w.r.t. the Chebyshev terms, denoted as  $\nabla_k \mathcal{L} = \frac{\partial \mathcal{L}}{\partial T_k(\mathbf{L})}$ , and since  $\mathbf{L}$  is shared across  $\{T_k(\mathbf{L})\}_k$ , one may write

$$\frac{\partial \mathcal{L}}{\partial \mathbf{L}} = \text{vec}^{-1} \left( \sum_{k=0}^{K-1} \mathbf{J}_k \text{vec}(\nabla_k \mathcal{L}) \right), \quad (2)$$

being  $\mathbf{J}_k \in \mathbb{R}^{n^2 \times n^2}$  the diagonal Jacobian matrix whose entry  $[\mathbf{J}_k]_{ij,ij} = \frac{\partial [T_k(\mathbf{L})]_{ij}}{\partial \mathbf{L}_{ij}}$  and  $\text{vec}(\cdot)$  a vectorization that appends the entries of a given matrix using the x-y order in  $\mathbf{J}_k$ , and  $\text{vec}^{-1}$  its inverse. In the above equation, one may show that each entry of the Jacobian  $\mathbf{J}_k$  can be recursively obtained as

$$\begin{cases} 0 & k = 0 \\ 1 & k = 1 \\ 2[[T_{k-1}(\mathbf{L})]_{ij} + \mathbf{L}_{ij} \frac{\partial [T_{k-1}(\mathbf{L})]_{ij}}{\partial \mathbf{L}_{ij}}] - \frac{\partial [T_{k-2}(\mathbf{L})]_{ij}}{\partial \mathbf{L}_{ij}} & k \geq 2, \end{cases} \quad (3)$$

so  $\mathbf{L}$  can be updated using Eqs (2), (3) and gradient descent. However, as designed above, the learned matrix  $\mathbf{L}$  is not guaranteed to be a valid Laplacian. In what follows, we consider reparametrization which constrains  $\mathbf{L}$  to be valid.

#### 3.1. Laplacian Reparametrization

We turn the design of  $\mathbf{L}$  into the learning of the adjacency matrix  $\mathbf{A}$  while guaranteeing the resulting matrix  $\mathbf{L}$  to be a valid Laplacian. Different reparametrizations are considered including  $\mathbf{L} = \mathbf{D}(\mathbf{A}) - \mathbf{A}$  which corresponds to the combinatorial form. If we constrain  $\mathbf{A}$  to be column-stochastic, then  $\mathbf{L}$  corresponds to the random walk graph Laplacian. We also consider other reparametrizations including the normalized defined as  $\mathbf{L} = \mathbf{I}_n - [\mathbf{D}(\mathbf{A}^{\top})]^{-\frac{1}{2}} \mathbf{A} [\mathbf{D}(\mathbf{A})]^{-\frac{1}{2}}$  (see table 1).

Constraints	Reparametrization	Jacobian
COMB	$\mathbf{D}(\mathbf{A}^\top) - \mathbf{A}$	$[\mathbf{J}_c]_{ij,pq} = 1_{\{i=j, p \neq q\}} - 1_{\{i \neq j\}}$
NDRW	$\mathbf{A} [\mathbf{D}(\mathbf{A})]^{-1}$	$[\mathbf{J}_{\text{ndrw}}]_{ij,pq} = 1_{\{j=q\}} (\delta_{ip} - \mathbf{L}_{ij}) [\mathbf{1}_n \mathbf{D}(\mathbf{A})^{-1}]_{pq}$
DRW	$\mathbf{I}_n - \mathbf{A} [\mathbf{D}(\mathbf{A})]^{-1}$	$[\mathbf{J}_{\text{drw}}]_{ij,pq} = 1_{\{j=q\}} (\mathbf{L}_{ij} - \delta_{ip}) [\mathbf{1}_n \mathbf{D}(\mathbf{A})^{-1}]_{pq}$
NDN	$[\mathbf{D}(\mathbf{A}^\top)]^{-\frac{1}{2}} \mathbf{A} [\mathbf{D}(\mathbf{A})]^{-\frac{1}{2}}$	$[\mathbf{J}_{\text{ndn}}]_{ij,pq} = 1_{\{i=p \vee j=q\}} \frac{\mathbf{L}_{ij}}{2\mathbf{A}_{pq}} (2\delta_{ip}\delta_{jq} - [\mathbf{D}(\mathbf{A}^\top)^{-1} \mathbf{A} + \mathbf{A} \mathbf{D}(\mathbf{A})^{-1}]_{pq})$
DN	$\mathbf{I}_n - [\mathbf{D}(\mathbf{A}^\top)]^{-\frac{1}{2}} \mathbf{A} [\mathbf{D}(\mathbf{A})]^{-\frac{1}{2}}$	$[\mathbf{J}_{\text{dn}}]_{ij,pq} = 1_{\{i=p \vee j=q\}} \frac{\mathbf{L}_{ij}}{2\mathbf{A}_{pq}} ([\mathbf{D}(\mathbf{A}^\top)^{-1} \mathbf{A} + \mathbf{A} \mathbf{D}(\mathbf{A})^{-1}]_{pq} - 2\delta_{ip}\delta_{jq})$
Symmetry	$\mathbf{A} + \mathbf{A}^\top$	$[\mathbf{J}_s]_{ij,pq} = 1_{\{(i=p, j=q) \vee (i=q, j=p)\}}$
S-COMB	$\mathbf{D}(\mathbf{A} + \mathbf{A}^\top) - (\mathbf{A} + \mathbf{A}^\top)$	$\mathbf{J}_{\text{sc}} = \mathbf{J}_c \mathbf{J}_s$
S-NDRW	$(\mathbf{A} + \mathbf{A}^\top) [\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-1}$	$\mathbf{J}_{\text{sndrw}} = \mathbf{J}_{\text{ndrw}} \mathbf{J}_s$
S-DRW	$\mathbf{I}_n - (\mathbf{A} + \mathbf{A}^\top) [\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-1}$	$\mathbf{J}_{\text{sdrw}} = \mathbf{J}_{\text{drw}} \mathbf{J}_s$
S-NDN	$[\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-\frac{1}{2}} (\mathbf{A} + \mathbf{A}^\top) [\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-\frac{1}{2}}$	$\mathbf{J}_{\text{sndn}} = \mathbf{J}_{\text{ndn}} \mathbf{J}_s$
S-DN	$\mathbf{I}_n - [\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-\frac{1}{2}} (\mathbf{A} + \mathbf{A}^\top) [\mathbf{D}(\mathbf{A} + \mathbf{A}^\top)]^{-\frac{1}{2}}$	$\mathbf{J}_{\text{sdn}} = \mathbf{J}_{\text{dn}} \mathbf{J}_s$

Table 1. Different reparametrizations and the underlying Jacobians. In this table, COMB stands for ‘‘Combinatorial’’ Laplacian, NDRW for ‘‘Non Differential Random Walk’’, DRW for ‘‘Differential Random Walk’’, NDN for ‘‘Non Differential Normalized’’ Laplacian, and DN for ‘‘Differential Normalized’’ one. The symmetric variants of these Laplacians are prefixed by ‘‘S’’.

For any reparametrization of  $\mathbf{L}$ , the chain rule leads to

$$\frac{\partial \mathcal{L}}{\partial \mathbf{A}} = \text{vec}^{-1} \left( \mathbf{J} \text{vec} \left( \frac{\partial \mathcal{L}}{\partial \mathbf{L}} \right) \right), \quad (4)$$

with  $\frac{\partial \mathcal{L}}{\partial \mathbf{L}}$  obtained from Eq. 2 and  $\mathbf{J}$  being a sparse Jacobian matrix whose entry  $[\mathbf{J}]_{ij,pq} = [\frac{\partial \mathbf{L}_{ij}}{\partial \mathbf{A}_{pq}}]_{ij,pq}$ ; this matrix is given in table 1 for different Laplacian settings including the combinatorial and random walk which respectively capture the differential and non-differential properties of node features. We also consider the differential random walk – as a combination of these two Laplacians – obtained by plugging the latter into the former. All these Laplacians are built upon either symmetric or non-symmetric matrices  $\mathbf{A}$ . Note that symmetry is obtained using weight sharing, i.e., by constraining the upper and the lower triangular parts of  $\mathbf{A}$  to share the same entries. This is guaranteed by considering a reparametrization as  $\mathbf{A} + \mathbf{A}^\top$  (with  $\mathbf{A}$  being now a free matrix) and by tying pairwise symmetric entries of the gradient  $\frac{\partial \mathcal{L}}{\partial \mathbf{A}}$ ; this is equivalently obtained by multiplying the original gradient  $\frac{\partial \mathcal{L}}{\partial \mathbf{A}}$  by the Jacobian  $[\mathbf{J}_s]_{ij,pq} = 1_{\{(i=p, j=q) \vee (i=q, j=p)\}}$  which is again extremely sparse and highly efficient to evaluate.

### 3.2. Orthogonality

Learning multiple matrices  $\{T_k(\mathbf{L})\}_{k=0}^{K-1}$  allows us to capture different graph topologies when achieving aggregation and convolution, and this enhances the discrimination power of the GCN representations without increasing the actual number of training parameters (as also shown later in experiments). However, if aggregation produces, for a given  $u \in \mathcal{V}$ , linearly dependent vectors  $\mathcal{X}_u = \{\sum_{u'} [T_k(\mathbf{L})]_{uu'} \psi(u')\}_k$ , then convolution will also generate linearly dependent representations with an overestimated number of training parameters in the null space of  $\mathcal{X}_u$ . Besides, matrices  $\{T_1(\mathbf{L}), \dots, T_K(\mathbf{L})\}$  used for aggregation, may also correspond to overlapping and

redundant neighborhoods.

Provided that  $\{\psi(u')\}_{u' \in \mathcal{N}_r(u)}$  are linearly independent, and  $K$  upper-bounded by  $\text{rank}(\{\psi(u')\}_{u' \in \mathcal{N}_r(u)}) \leq \min(|\mathcal{V}|, s)$ , the condition that makes vectors in  $\mathcal{X}_u$  linearly independent reduces to orthogonality, i.e.,  $\langle T_k(\mathbf{L}), T_{k'}(\mathbf{L}) \rangle_F = 0, \forall k \neq k'$ , with  $\langle \cdot, \cdot \rangle_F$  being the Hilbert-Schmidt (or Frobenius) inner product defined as  $\langle T_k(\mathbf{L}), T_{k'}(\mathbf{L}) \rangle_F = \text{tr}(T_k(\mathbf{L})^\top T_{k'}(\mathbf{L}))$  with  $\text{tr}(\cdot)$  being the matrix trace operator. A sufficient condition that guarantees the orthogonality of the Chebyshev basis consists in taking the Laplacian  $2(\mathbf{L} - \lambda_{\min} \mathbf{I}_n) / (\lambda_{\max} - \lambda_{\min}) - \mathbf{I}_n$  instead of  $\mathbf{L}$  with  $\lambda_{\min}$  (resp.  $\lambda_{\max}$ ) being the smallest (resp. largest) eigenvalue of  $\mathbf{L}$ , and this guarantees the eigenvalues of the resulting matrix to be in  $[-1, +1]$  and also the minimality of  $\{T_k(\mathbf{L})\}_k$  (see for instance [47]). It is easy to see that this normalization equates the rescaled Laplacian shown in section 2, i.e., on the combinatorial setting, as its smallest eigenvalue is zero.

## 4. Experiments

In this section, we evaluate the performance of our GCN network for the task of action recognition [93, 99] using two challenging skeleton datasets; SBU Interaction [88] and First-Person Hand Action (FPHA) [87]. The purpose is to show the relevance of our Laplacian design and its comparison against different handcrafted Laplacians and learned ones as well as more general related work in action recognition.

### 4.1. Datasets and implementation details

**Dataset description.** SBU is an interaction dataset acquired (under relatively well controlled conditions) using the Microsoft Kinect sensor; it includes in total 282 moving skeleton sequences (performed by two interacting individuals) belonging to 8 categories: ‘‘approaching’’, ‘‘departing’’, ‘‘pushing’’, ‘‘kicking’’, ‘‘punching’’, ‘‘exchanging objects’’,

“hugging”, and “hand shaking”. Each pair of interacting individuals corresponds to two 15 joint skeletons and each joint is encoded with a sequence of its 3D coordinates across video frames. In this dataset, we consider the same evaluation protocol as the one suggested in the original dataset release [88] (i.e., train-test split).

The FPHA dataset includes 1175 skeletons belonging to 45 action categories which are performed by 6 different individuals in 3 scenarios. In contrast to SBU, action categories are highly variable with inter and intra subject variability including style, speed, scale and viewpoint. Each skeleton includes 21 hand joints and each joint is again encoded with a sequence of its 3D coordinates across video frames. We evaluate the performance of our method using the 1:1 setting proposed in [87] with 600 action sequences for training and 575 for testing. In all these experiments, we report the average accuracy over all the classes of actions.

**Skeleton normalization.** Let  $S^t = \{p_1^t, \dots, p_n^t\}$  denote the 3D skeleton coordinates at frame  $t$ . Without a loss of generality, we consider a particular order so that  $p_1^t$ ,  $p_2^t$  and  $p_3^t$  correspond to three reference joints (e.g., neck, left shoulder and right shoulder for SBU dataset); as shown in Fig. 3, this corresponds to joints 2, 4 and 7 for SBU and 1, 3 and 5 for FPHA. As the relative distance between these 3 joints is stable w.r.t. any motion, these 3 joints are used in order to estimate the rigid motion (similarity transformation) for skeleton normalization (see also [89]). Each graph sequence is processed in order to normalize its 3D coordinates using a similarity transformation; the translation parameters  $\mathbf{t} = (t_x, t_y, t_z)$  of this transformation correspond to the shift that makes the reference point  $(p_2^0 + p_3^0)/2$  coincide with the origin while the rotation parameters  $(\theta_x, \theta_y, \theta_z)$  are chosen in order to make the plane formed by  $p_1^0$ ,  $p_2^0$  and  $p_3^0$  coplanar with the x-y plane and the vector  $p_2^0 - p_3^0$  colinear with the x-axis. Finally, the scaling  $\gamma$  of this similarity is chosen to make the  $\|p_2^0 - p_3^0\|_2$  constant through all the action instances. Hence, each normalized joint is transformed as  $\hat{p}_i^t = \gamma(p_i^t - \mathbf{t})R_x(\theta_x)R_y(\theta_y)R_z(\theta_z)$  with  $R_x$ ,  $R_y$ ,  $R_z$  being rotation matrices along the  $x$ ,  $y$  and  $z$  axes respectively.

**Input graphs.** Considering a sequence of normalized skeletons  $\{S^t\}_t$ , each joint sequence  $\{\hat{p}_j^t\}_t$  in these skeletons defines a labeled trajectory through successive frames (see Fig. 2). Given a finite collection of trajectories, we consider the input graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$  where each node  $v_j \in \mathcal{V}$  corresponds to the labeled trajectory  $\{\hat{p}_j^t\}_t$  and an edge  $(v_j, v_i) \in \mathcal{E}$  exists between two nodes iff the underlying trajectories are spatially neighbors. Each trajectory (i.e., node in  $\mathcal{G}$ ) is processed using *temporal chunking*: first, the total duration of a sequence (video)

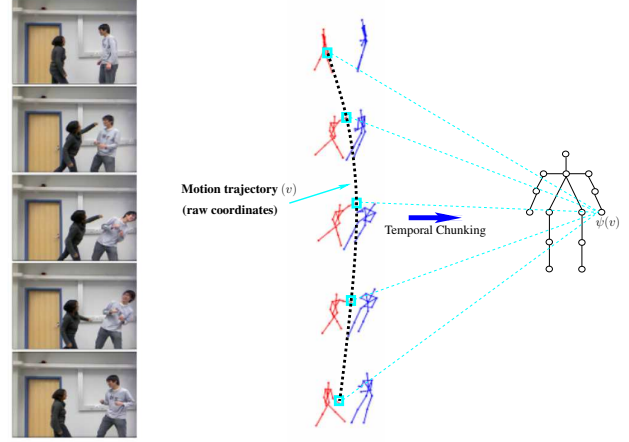


Figure 2. This figure shows the whole keypoint tracking and description process.

is split into  $M$  equally-sized temporal chunks ( $M = 4$  in practice), then the normalized joint coordinates  $\{\hat{p}_j^t\}_t$  of the trajectory  $v_j$  are assigned to the  $M$  chunks (depending on their time stamps) prior to concatenate the averages of these chunks; this produces the description of  $v_j$  (again denoted as  $\psi(v_j) \in \mathbb{R}^s$  with  $s = 3 \times M$ ) and  $\{\psi(v_j)\}_j$  constitutes the raw description of nodes in a given sequence. Note that two trajectories  $v_j$  and  $v_i$ , with similar joint coordinates but arranged differently in time, will be considered as very different when using temporal chunking. Note also that beside being compact and discriminant, this temporal chunking gathers advantages – while discarding drawbacks – of two widely used families of techniques mainly *global averaging techniques* (invariant but less discriminant) and *frame resampling techniques* (discriminant but less invariant). Put differently, temporal chunking produces discriminant raw descriptions that preserve the temporal structure of trajectories while being *frame-rate* and *duration* agnostic.

**Implementation settings.** We trained the GCN networks end-to-end using the Adam optimizer [101] for 1,800 epochs with a batch size equal to 200 for SBU and 600 for FPHA, a momentum of 0.9 and a global learning rate (denoted as  $\nu(t)$ ) inversely proportional to the speed of change of the loss used to train our networks; when this speed increases (resp. decreases),  $\nu(t)$  decreases as  $\nu(t) \leftarrow \nu(t-1) \times 0.99$  (resp. increases as  $\nu(t) \leftarrow \nu(t-1)/0.99$ ). All these experiments are run on a GeForce GTX 1070 GPU device (with 8 GB memory) and neither dropout nor data augmentation are used.

## 4.2. Baselines

We compare the performances of our GCN w.r.t. different Chebyshev basis settings which are defined upon handcrafted and learned Laplacians, as well as against totally learned Laplacian basis. Note that all these Laplacians are combined with symmetry and orthogonality settings as described earlier.

**Handcrafted Laplacians (HL).** All the Chebyshev terms  $\{T_k(\cdot)\}_k$  are evaluated upon a *handcrafted* Laplacian  $\mathbf{L}$  which in turns depends on a fixed adjacency matrix  $\mathbf{A}$  (set using the original input graph).

**Multi-Laplacians (ML).** In this configuration, the Laplacian used in the Chebyshev terms is trained as a weighted *combination* of the handcrafted variants of the Laplacians in table 1 (built upon the fixed matrix  $\mathbf{A}$ ). Note that orthogonality is obtained by normalizing the final learned Laplacian while symmetry is enforced in the handcrafted adjacency matrix  $\mathbf{A}$ .

**Totally Learned Laplacians (TLL).** In this variant,  $K$  independent Laplacians  $\{\mathbf{L}_k\}_k$  (and hence the underlying adjacency matrices) are learned. In contrast to the handcrafted setting, orthogonality and symmetry are obtained as a part of the optimization process (as already discussed in sections 3.1 and 3.2).

Laplacians Settings		Differential COMB	Non-Differential NDRW NDN		Combined DRW DN	
$K = 2$	HL	96.9231	96.9231	96.9230	93.8462	96.9230
	TLL	96.9231	95.3846	98.4615	96.9231	98.4615
	Our	98.4615	98.4615	96.9230	96.9231	98.4615
$K = 4$	HL	95.3846	93.8462	96.9231	96.9230	96.9230
	TLL	96.9231	95.3846	98.4615	98.4615	96.9230
	Our	98.4615	<b>100.000</b>	98.4615	98.4615	98.4615
$K = 8$	HL	96.9231	98.4615	96.9231	96.9230	96.9230
	TLL	96.9231	98.4615	98.4615	93.8462	96.9230
	Our	96.9231	98.4615	98.4615	98.4615	98.4615

Table 2. Detailed performances on SBU using Chebyshev networks with handcrafted (HL) and learned Laplacians (Our), and using totally learned Laplacians (TLL). These performances are shown for  $K \in \{2, 4, 8\}$  and for different reparametrizations of the Laplacians including differential (COMB), and non differential (NDRW, NDN) as well as their combinations (DRW, DN); see again Table. 1. Note that both symmetry and orthogonality constraints are used in these results.

## 4.3. Ablation study and comparison

Tables 2, 3 show a comparison of our GCN-based action recognition against the aforementioned GCN baselines, i.e. based on Handcrafted Laplacians and Totally Learned ones (performances with Multi-Laplacians are rather shown in Table 5); these comparisons are shown for different  $K \in \{2, 4, 8\}$ . From all these results, we observe a clear

Laplacians Settings		Differential COMB	Non-Differential NDRW NDN		Combined DRW DN	
$K = 2$	HL	85.9130	85.3913	85.5652	85.3913	84.8695
	TLL	85.5652	86.4348	85.7391	85.3913	86.0869
	Our	85.3913	85.7391	85.5652	85.5652	85.7391
$K = 4$	HL	86.4348	84.1739	85.9130	84.0000	84.5217
	TLL	84.3478	85.3913	86.4347	85.0435	85.5652
	Our	85.2174	85.3913	85.7391	<b>87.1304</b>	<b>87.3043</b>
$K = 8$	HL	85.2174	83.8261	86.0869	84.6957	85.7391
	TLL	84.5217	85.5652	85.7391	85.0435	84.8695
	Our	84.6957	86.9565	86.7826	84.6957	84.5217

Table 3. Same caption as Table. 2 on the FPHA database.

gain of our Chebyshev-based Laplacian design w.r.t. these baselines; at least one of the setting (namely  $K = 4$ ) provides a significant gain. Table 4 shows an ablation study, where the impact of each component of our GCN (Laplacians, symmetry and orthogonality) is observed separately and jointly. From these results, we observe a positive impact when constraining the learned matrices to be symmetric and orthogonal; this gain is noticeable with non-differential Laplacians on SBU and with combined (differential/non-differential) ones on FPHA and this clearly shows the complementary aspect of these two Laplacian settings mainly on challenging datasets (i.e., FPHA). Again, this gain reaches the highest values when  $K$  is sufficiently (not very) large and this follows the small size of the original skeletons (diameter and dimensionality of the graphs and the signal) used for action recognition which constrains the required number of Laplacian terms in the Chebyshev decomposition. Hence, with few Chebyshev terms, our method is able to learn relevant Laplacians and representations for action recognition.

Dataset	Constraints Sym Orth		Differential COMB	Non-Differential NDRW DRW		Combined NDN DN		Avg. perf.
	Sym	Orth						
SBU	$\times$	$\times$	90.76	98.46	98.46	96.92	98.46	96.61
	$\checkmark$	$\times$	95.38	100.0	98.46	98.46	98.46	98.15
	$\times$	$\checkmark$	95.38	96.92	98.46	96.92	98.46	97.23
	$\checkmark$	$\checkmark$	98.46	<b>100.00</b>	98.46	98.46	98.46	<b>98.76</b>
Avg.	-	-	95.00	<b>98.84</b>	98.46	97.69	98.46	-
FPHA	$\times$	$\times$	79.30	85.21	86.60	86.78	86.43	84.86
	$\checkmark$	$\times$	84.00	85.73	85.73	86.43	86.08	85.60
	$\times$	$\checkmark$	83.13	85.39	85.73	86.78	86.78	85.56
	$\checkmark$	$\checkmark$	85.21	85.39	85.73	<b>87.13</b>	<b>87.30</b>	<b>86.15</b>
Avg.	-	-	82.91	85.43	85.95	<b>86.78</b>	<b>86.65</b>	-

Table 4. Ablation study on SBU and FPHA databases, when symmetry (sym) and orthogonality (orth) are taken separately and when combined (in these results  $K = 4$ ).

Our proposed Laplacian design avoids the strong bias about the handcrafted adjacency matrices which are rather suitable to capture the anthropometric characteristics of skeletons and less optimal for action recognition. On another hand, Tables. 2, 3 and 4 show that our Laplacian design makes it possible to capture better the topology of the graph data (i.e., the neighborhood system defined by the learned Laplacian and its underlying adjacency matrix



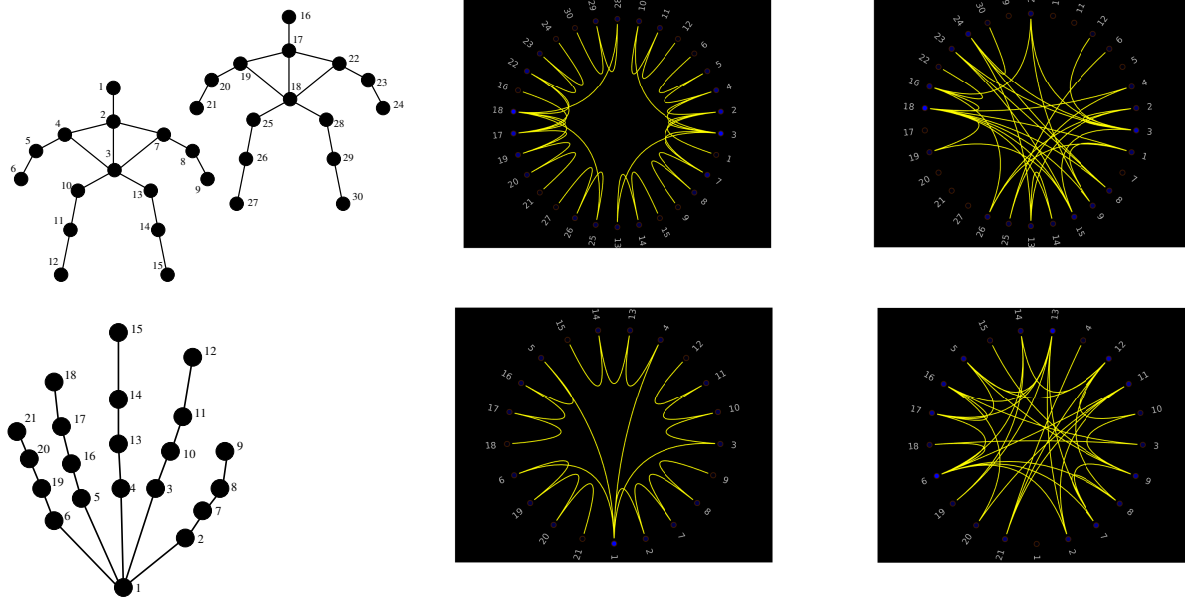


Figure 3. This figure shows original skeletons (left) with their intrinsic node-to-node relationships useful for *individual identification* (middle), and an example of the adjacency matrix associated to the learned Laplacian which shows the extrinsic node-to-node relationships found to be the most discriminating for *skeleton-based action recognition* when using our proposed method (the exact setting corresponds to Tables 2 and 3, using NDRW Laplacian with  $K = 4$ ). (Better to zoom the PDF version to view the learned node-to-node relationships).

A). In contrast, the baselines are limited when connectivity is handcrafted and also when learned using totally trained Laplacians, as this results either into a biased Laplacian or into a larger number of training parameters, while Chebyshev provides a compromise between these two extreme cases: indeed, it enhances the discrimination power of the representation without increasing the actual number of training parameters. In sum, the gain of our GCN results from (i) the relative flexibility of the proposed design which allows learning complementary aspects of graph topology (through the Chebyshev basis), and also (ii) the regularization effect of our constraints (Laplacian weight sharing in Chebyshev, Laplacian reparametrization, orthogonality and symmetry) which all mitigate overfitting.

Finally, we compare the classification performances of our GCN against other, and more general, related methods in action recognition ranging from sequence based such as LSTM and GRU [18, 90, 104] to deep graph (non-vectorial) methods [24], etc. (see related work in tables 5 and 6). From the results in these tables, our GCN brings a noticeable gain w.r.t. related state of the art methods.

## 5. Conclusion

We introduce in this paper a novel Chebyshev-based Laplacian design for graph convolutional networks (GCNs). The learned Laplacians capture the most influencing interactions between body parts in skeleton action recognition. The strength of our method resides in its ability to

Method	Accuracy (%)
Raw Position [88]	49.7
Joint feature [32]	86.9
CHARM [33]	86.9
H-RNN [16]	80.4
ST-LSTM [17]	88.6
Co-occurrence-LSTM [30]	90.4
STA-LSTM [18]	91.5
ST-LSTM + Trust Gate [17]	93.3
VA-LSTM [19]	97.6
GCA-LSTM [104]	94.9
Riemannian manifold. traj [103]	93.7
DeepGRU [90]	95.7
RHCN + ACSC + STUFE [24]	98.7
Multi-Laplacians (ML baseline [61])	98.4
Our best (table 2)	<b>100</b>

Table 5. Comparison against state of the art methods using the SBU database.

learn shared Laplacians which are embedded in a Chebyshev basis that increases the discrimination power of our graph representation. In contrast to usual deep networks, the parameters of our GCNs are interpretable as their design is constrained “by construction” and this also brings a regularization effect that mitigates overfitting. Indeed, our Laplacian weight sharing and reparametrization — together with symmetry and orthogonality constraints — enhance the representational power of our learned graph features without increasing the actual number of training parameters in the resulting GCNs. Several operators are considered in-



Method	Color	Depth	Pose	Accuracy (%)
Two stream-color [91]	✓	✗	✗	61.56
Two stream-flow [91]	✓	✗	✗	69.91
Two stream-all [91]	✓	✗	✗	75.30
HOG2-depth [92]	✗	✓	✗	59.83
HOG2-depth+pose [92]	✗	✓	✓	66.78
HON4D [94]	✗	✓	✗	70.61
Novel View [95]	✗	✓	✗	69.21
1-layer LSTM [30]	✗	✗	✓	78.73
2-layer LSTM [30]	✗	✗	✓	80.14
Moving Pose [96]	✗	✗	✓	56.34
Lie Group [23]	✗	✗	✓	82.69
HBRNN [16]	✗	✗	✓	77.40
Gram Matrix [97]	✗	✗	✓	85.39
TF [98]	✗	✗	✓	80.69
JOULE-color [100]	✓	✗	✗	66.78
JOULE-depth [100]	✗	✓	✗	60.17
JOULE-pose [100]	✗	✗	✓	74.60
JOULE-all [100]	✓	✓	✓	78.78
Huang et al. [13]	✗	✗	✓	84.35
Huang et al. [43]	✗	✗	✓	77.57
Our best (table 3)	✗	✗	✓	<b>87.3</b>

Table 6. Comparison against state of the art methods using the FPHA database.

cluding differential and non-differential Laplacians which model the statistical properties of the learned graph representations, and when combined, they further enhance the performances of action recognition. Extensive experiments, conducted on standard databases (namely SBU and FPHA), show a clear gain of our design w.r.t. different handcrafted and (other) learned Laplacians, as well as the related work in skeleton-based action recognition.

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