Unsupervised classification of SDSS galaxy spectra

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ABSTRACT

Context. Defining templates of galaxy spectra is useful to quickly characterise new observations and organise databases from surveys. These templates are usually built from a pre-defined classification based on other criteria. This results in a somewhat complicated scheme with 38 classes (Dobos et al. 2012) or in a rather general classification with only 6 classes (Wang et al. 2018). More in-depth classifications of spectra are driven by several spectral features such as emission lines or their combinations with colours, nuclear, and star formation activity and morphology. This results in a somewhat complicated scheme with 38 classes (Dobos et al. 2012) or in a rather general classification with only 6 classes (Wang et al. 2018).

Methods. The spectra were first corrected for redshift, then wavelet-filtered to reduce the noise, and finally binned to obtain about 1437 wavelengths per spectrum. The unsupervised clustering algorithm Fisher-EM, relying on a discriminative latent mixture model, was applied on these corrected spectra. The full set and several subsets of 100,000 and 300,000 spectra were analysed.

Results. The optimum number of classes given by a penalised likelihood criterion is 86 classes, of which the 37 most populated gather 99% of the sample. These classes are established from a subset of 302,214 spectra. Using several cross-validation techniques we find that this classification agrees with the results obtained on the other subsets with an average misclassification error of about 15%. The large number of very small classes tends to increase this error rate. In this paper, we do an initial quick comparison of our classes with literature templates.

Conclusions. This is the first time that an automatic, objective and robust unsupervised classification is established on such a large number of galaxy spectra. The mean spectra of the classes can be used as templates for a large majority of galaxies in our Universe.

Key words. methods: data analysis – methods: statistical – galaxies: statistics – galaxies: general – techniques: spectroscopic

1. Introduction

There is a long recognised need for templates of typical galaxy spectra to quickly characterise new objects. For instance, such templates are used in automatic classification in databases such as the Sloan Digital Sky Survey (SDSS), in which a χ² minimisation is used to separate galaxies from quasars (QSOs) and stars, followed by a categorisation into starbursts, star-forming galaxies and active galactic nuclei (AGN) using line ratios.¹

Templates can be built by stacking spectra to reduce the noise that can make some features disappear (Dobos et al. 2012). This strategy obviously requires perfectly identical spectra that can be obtained by visual inspection, by a classification made from some general or specific properties of the galaxies (e.g., line ratios, continuum, morphology, and model fitting of stellar population), or by machine learning tools.

Kennicutt (1992) proposed an atlas of spectra classified according to the morphology of the galaxies. This kind of approach makes the assumption of a perfect correspondence between the two pieces of information and is insufficient to characterise the informative wealth of the spectra.

¹ The mean spectra of the 86 classes and the class of each of the 302,248 spectra are only available at the CDS via anonymous ftp to cdsarc.u-strasbg.fr (130.79.128.5) or via http://cdsarc.u-strasbg.fr/viz-bin/cat/J/A+A/649/A53
² https://www.sdss.org/dr12/spectro/catalogs/
#Objectinformation
the classes. Wang et al. (2018) then compare the median com-
posite spectra themselves simply using the a priori correspond-
ence of the classes. They conclude that the continua match well
but not the emission lines.

The main drawback of most of these studies is that they
mainly rely on a visual inspection. This limits the sample size
used to establish the templates and thus their representativ-
ness of the galaxy diversity. New methods should make use of
the information contained in the spectra themselves to become
both automatic and objective, and machine learning tools of
unsupervised classification (clustering) are well suited for this
(Fraix-Burnet et al. 2015).

The goal of unsupervised classification is to detect structures
in the data space from which clusters of similar spectra can be
identified. The average spectrum of a cluster defines a typical
template that is dictated only by the data. The physics comes
afterwards, by inverse procedures or model fitting to charac-
terise the underlying stellar population and the ionisation con-
ditions within the corresponding galaxy (e.g., Moulkata et al.
2004; Noll et al. 2009; Comparat et al. 2017). Then, these clus-
ters can be considered as classes.

Principal component analysis (PCA) defines an orthogonal
base of spectra from which templates can be constructed. For
instance, Marchetti et al. (2013) find that three principal compo-
nents are sufficient to describe the variance of their sample, and
the coefficients (loadings) of each spectra are used to perform an
unsupervised k-means classification. However, the variance axes
are not necessarily the discriminant axes (Chang 1983).

Other ways of classifying spectra are to use the embedded
information, such as the independent component analysis (ICA,
Lu et al. 2006) or t-distributed stochastic neighbour embedding
(t-SNE) reduction of spectral information which, to our knowl-
edge, has only been applied on stellar spectra (Traven et al.
2017). Some efficient tools can be designed for specific pur-
poses, such as the detection of outliers (Baron & Poznanski
2017).

So far, no robust unsupervised classification of galaxy spec-
tra has been proposed. An attempt with raw spectra and the
k-means approach was performed by Sánchez Almeida et al.
(2010) but this result is disputed by De et al. (2016). Indeed, this
technique does not perform well in high dimension, when the
classes are not balanced and well separated in parameter space.

A more powerful tool is the Gaussian mixture model (GMM,
Bouveyron et al. 2019) approach which fits a multivariate
Gaussian to each cluster (e.g., de Souza et al. 2017). Mixture
models are particularly popular in the context of clustering since
they facilitate fitting to various situations and to automatically
find the appropriate number of clusters in the data at hand. In
addition, in the case of a high-dimensional space, subspace clus-
tering usually yields more robust results that can be both justified
from the statistical point of view (curse of dimensionality) and
the physical argument that not all parameters are discriminant.
This is especially true for spectra because of the noise and many
uninformative wavelengths.

In this paper, we use a GMM approach implemented in the
Fisher-EM algorithm (Bouveyron & Brunet 2012a). It is a sub-
space clustering algorithm, based on a latent mixture model
that fits the data into a low-dimensional discriminative sub-
space. Fisher-EM uses a modified version of the expectation–
maximisation (EM) algorithm for model inference and intro-
duces a Fisher criterion to optimise the quality of the clustering.
It has been used only once in astrophysics up to now, on a data
set of 100 000 galaxies described by 12 parameters (Siudek et al.
2018). In the present paper, we apply Fisher-EM on a larger
sample of 702 248 spectra (with 1437 wavelengths) from the
SDSS.

The paper is organised as follows. The data are described in
Sect. 2. The Fisher-EM algorithm and its application on our large
data set are detailed in Sect. 3. The results are given in Sect. 4
and an estimation of the statistical robustness of our classification
is presented in Sect. 6. To get a first idea of the physical meaning
of our classes, we present a comparison with the existing atlases of
Kennicutt (1992), Dobos et al. (2012), Wang et al. (2018) in
Sect. 7. A discussion and a conclusion are given in Sects. 7 and 8
respectively.

2. The data

In this work, we use the same data set as studied by
Sánchez Almeida et al. (2010) and De et al. (2016). Thus, we
retrieved the spectra of 702 248 galaxies and QSOs (with red-
shift smaller than 0.25) from the SDSS database, release 7. For
consistency with the two previous studies, we chose not to cor-
rect the spectra for galactic extinction. Apart from a philosophi-
ical justification explained in Sánchez Almeida et al. (2010), the
correction would be small (median extinction $g−r$ of 0.03 for
our sample) and unavoidably somewhat uncertain, bringing an
additional variance (see a further comment in Sect. 7.1). The raw
spectra have 3850 points in the wavelength range imposed by the
instrument, $\lambda = 3800$–9250 Å. The spacing is uniform in reso-
lation ($\delta \lambda/\lambda = 1/4342$). Taking the redshift into account, the
range common to all the spectra goes from 3806 to 7371 Å with
2874 wavelengths. We then corrected the spectra for the redshift,
as in De et al. (2016), using the Shannon criterion to preserve
the shape of the spectral lines. For this purpose, we doubled the
sampling of the spectra beforehand, generating 5748 points for
each spectrum. The resulting spectra were then normalised by
dividing each spectrum by its average value between 4300 and
5000 Å.

Contrary to Sánchez Almeida et al. (2010) and De et al.
(2016), we did not select a priori bands supposedly containing
relevant physical information to reduce the size of data array.
Instead, we first applied a noise reduction through wavelet fil-
tering followed by a binning by a factor of 4 to obtain spectra
with 1437 wavelengths, which is a number comparable with that
used in the two previous works. In this process, we keep most of
the information with the shapes of the lines being preserved
(see Appendix A) as well as full objectivity. These computations
were done using algorithms under the R environment3.

3. The Fisher-EM algorithm

3.1. Mathematical overview of the Fisher-EM algorithm

The Fisher-EM algorithm relies on a statistical model, named
the discriminative latent mixture (DLM) model, and uses a modified
version of the EM algorithm for model inference. We provide
hereafter a short mathematical overview of both the model and
the inference algorithm.

The DLM model assumes that $[y_1, \ldots, y_n] \in \mathbb{R}^P$ denotes the
data set of $n$ observations that we want to cluster into $K$ homo-
geneous groups, that is adjacent to each observation $y_i$ a value
$z_i \in \{1, \ldots, K\}$ where $z_i = k$ indicates that the observation $y_i$
belongs to the $k$th group. On the one hand, let us assume that

\footnote{2}{http://www.sdss.org/dr7/}
\footnote{3}{https://www.r-project.org/}, algorithms \texttt{wd}, \texttt{threshold} and \texttt{wr}
from the package \texttt{wavethresh}, and \texttt{binning} from the package \texttt{prospectr}.

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$[y_1, \ldots, y_n]$ are independent observed realisations of a random vector $Y \in \mathbb{R}^p$ and that $[z_1, \ldots, z_n]$ are also independent realisations of a random variable $Z \in \{1, \ldots, K\}$. On the other hand, let $\mathbb{E} \subset \mathbb{R}^p$ denote a latent space assumed to be the most discriminative subspace of dimension $d \leq K - 1$ such that $0 \in \mathbb{E}$. Moreover, let $[x_1, \ldots, x_n] \in \mathbb{E}$ denote the actual data, described in the latent space $\mathbb{E}$ of dimension $d$, which are in addition presumed to be independent realisations of an unobserved random vector $X \in \mathbb{E}$. Finally, the observed variable $Y \in \mathbb{R}^p$ and the latent variable $X \in \mathbb{E}$ are assumed to be linked through a linear transformation as follows:

$$Y = UX + \varepsilon,$$

(1)

where $U$ is a $p \times d$ orthonormal matrix common to the $K$ groups and satisfying $U^TU = I_d$. The $p$-dimensional random vector $\varepsilon$ stands for the noise term that models the non-discriminative information and is assumed to be distributed according to a centred Gaussian density function with a covariance matrix $\Psi$ ($\varepsilon_k \sim N(0, \Psi_k)$). Besides, within the latent space, $X$ is assumed, conditionally to $Z = k$, to be Gaussian, where

$$X_{ik|Z=k} \sim N(\mu_k, \Sigma_k)$$

(2)

and $\mu_k \in \mathbb{R}^d$ and $\Sigma_k \in \mathbb{R}^{d \times d}$ are the mean vector and the covariance matrix of the $k$th group, respectively. Given these distribution assumptions and according to Eq. (1),

$$Y_{[X,Z=k]} \sim N(UX, \Psi_k),$$

(3)

and its marginal distribution is therefore a mixture of Gaussians as follows:

$$f(y) = \sum_{k=1}^{K} \pi_k \phi(y; m_k, S_k),$$

(4)

where $\pi_k$ is the mixing proportion of the $k$th group and $\phi(\cdot; m_k, S_k)$ denotes the multivariate Gaussian density function parametrised by the mean vector $m_k = U\mu_k$ and the covariance matrix $S_k = US_kU^T + \Psi_k$ of the $k$th group. Furthermore, we define the $p \times p$ matrix $W = [U, V]$ such that $W^TW = WW^T = I_p$, where the $(p - d) \times p$ matrix $V$ is an orthogonal complement of $U$. Finally, the noise covariance matrix $\Sigma_k$ is assumed to satisfy the conditions $V^T\Sigma_kV = \beta_kI_{p-d}$ and $U^T\Sigma_kU = 0_d$, such that $\Delta_k = W^T\Sigma_kW$ has the following form:

$$\Delta_k = \begin{pmatrix} \Sigma_k & \beta_k \beta_k^T & \beta_k^T \beta_k \\ \beta_k & \beta_k & \beta_k \\ 0 & \beta_k & \beta_k \end{pmatrix},$$

$$d \leq K - 1,$$

$$p - d$$

These last conditions imply that the discriminative and the non-discriminative subspaces are orthogonal, which suggests in practice that all the relevant clustering information remains in the latent subspace. From a practical point of view, $\beta_k$ models the variance of the non-discriminative noise of the data. The whole model is referred to by $DLM_{[\Sigma_k, \beta_k]}$ in Bouveyron & Brunet (2012a).

Several other models can be obtained from the $DLM_{[\Sigma_k, \beta_k]}$ model by relaxing or adding constraints on model parameters. For instance, the covariance matrices $\Sigma_1 \ldots \Sigma_K$ and the noise parameters $\beta_1 \ldots \beta_K$ can be assumed to be common to all classes (hence independent on $k$). This is the statistical model $DLM_{[\Sigma_0]}$ (referred to as DB in the R package FisherEM). A list of the 12 different DLM models is given in Table 1 in Bouveyron & Brunet (2012a). Such a family simultaneously yields very parsimonious models and allows us to carry out a fit in various situations.

An estimation procedure, called the Fisher-EM algorithm, is also proposed in Bouveyron & Brunet (2012a) to estimate the discriminative space and parameters of the mixture model. This algorithm is based on the EM algorithm to which an additional step is introduced between the E and M-step. This additional step, named F-step, aims to compute the projection matrix $U$ whose columns span the discriminative latent space. The Fisher-EM algorithm has therefore the following form, at iteration $q$:

- E-step: This step computes the posterior probabilities $t_{ik}^{(q)}$ that the observations belong to the $k$ groups using the following update formula:

$$t_{ik}^{(q)} = \hat{\pi}_k \hat{\beta}_k \phi(y_i; \hat{\theta}_k^{(q-1)}),$$

(5)

where $\hat{\theta}_k = [\hat{\mu}_k, \hat{\Sigma}_k, \hat{\beta}_k, \hat{U}]$.

- F-step: This step estimates, conditionally to the posterior probabilities, the orientation matrix $U^{(q)}$ of the discriminative latent space by maximising Fisher’s criterion (defined by the ratio of the between-class variance to the within-class variance, Fisher 1936; Fukunaga 2013) under the following orthonormality constraints:

$$\hat{U}^{(q)} = \max_{U'} \text{trace} \left( (U'SU)^{-1} U'S^{(q)} U' \right),$$

w.r.t. $U'U = I_d$,

(6)

where $S$ stands for the covariance matrix of the whole data set and $S^{(q)}_B$ defined as follows:

$$S^{(q)}_B = \frac{1}{n} \sum_{k=1}^{K} n_k^{(q)} (m_k^{(q)} - \bar{y}) (m_k^{(q)} - \bar{y})',$$

(7)

denotes the soft between-class covariance matrix with $n_k^{(q)} = \sum_{i=1}^{n_k} t_{ik}^{(q)}$, $m_k^{(q)} = \sum_{i=1}^{n_k} y_i / n_k^{(q)}$ and $\bar{y} = \sum_{i=1}^{n} y_i / n$.

- M-step: This third step estimates the parameters of the mixture model in the latent subspace by maximising the conditional expectation of the complete log-likelihood.

The Fisher-EM procedure iteratively updates the posterior probabilities and the parameters until the Aitken criterion is satisfied (see Paragraph 4.5 of Bouveyron & Brunet 2012a). The convergence properties of the Fisher-EM algorithm were studied in Bouveyron & Brunet (2012b). Finally, since the latent subspace has a low dimension and is common to all groups, the clustered data can be easily visualised by projecting them into the estimated latent subspace.

The choice of the best statistical DLM model and the optimum number of clusters depends on the data and are estimated with three likelihood criteria: the classical Akaike information criterion (AIC), the popular Bayesian information criterion (BIC) and the integrated completed likelihood (ICL). These criteria all penalise the likelihood by the number of parameters of the statistical model, but the BIC also includes the number of observations while the ICL in addition favours well separated statistical models (Biernacki et al. 2000; Girard & Saracco 2016). Hence, the maximum value of ICL determines the best statistical model and the optimum number $K$ of classes in the analyses performed in the present study.
The Fisher-EM algorithm is implemented in the eponym package for R. For the present study, we used the version 1.5.1 of this package.

### 3.2. Differences between k-means and Fisher-EM

The clustering algorithm k-means is relatively standard and easy to use. However, this algorithm has severe limitations that triggered the development of generalisations, Fisher-EM being one of the most recent and complete. The statistical differences can be summarised as follows:

1. **K-means** assumes that all classes can be fit with the same pattern, that is a spherical Gaussian. The Fisher-EM algorithm belongs to the class of GMMs that releases this strong assumption.

2. **K-means** is a hard clustering algorithm: each observation belongs to one class and only one. Fisher-EM, which is based on the EM algorithm, is soft in the sense that each observation is associated with a vector of probabilities for all classes. In Fisher-EM, an observation is attributed to a class if the corresponding probability is higher than 0.5.

3. **K-means** works in the full data space, while Fisher-EM looks for a subspace in which the classes are more conspicuous, and thus makes a selection of the most pertinent dimensions.

4. **K-means** uses only the Euclidean distance to define classes, while Fisher-EM takes into account the variance and optimises the compactness of classes.

5. **K-means** always yields a solution, while Fisher-EM, as all EM algorithms, may not be able to converge to a local maximum (Wu 1983; Bouveyron & Brunet 2012b).

All possible classes of galaxy spectra can certainly not be represented by either spherical or identical distributions in the data space, and many wavelengths bring either no information or just noise. These two points favour the use of subspace and GMM-based clustering algorithms for large samples of astrophysical spectra.

### 3.3. Application to the large data set of SDSS spectra

The selection of the best statistical model and the optimum number of classes requires many computations. Since our data set is very large, it is inconvenient in practice to use the full sample at once. With a low number of classes, say \( K = 3 \), the analysis with Fisher-EM takes about 30 h CPU time for 702 248 spectra. But for \( K = 50 \), on a smaller sample of 300 000 spectra, it takes several hundreds of hours. Considering that the computation has to be repeated for each of the 12 statistical models and for many values of \( K \) to find the maximum ICL value, splitting the sample into subsets makes the analyses manageable and allows for robustness estimation.

We thus divided the full sample into several subsets that have been analysed separately and for different purposes. We are aware that this approach results in a somewhat complicated scheme, but in this exploring stage of the application of Fisher-EM on such a large astronomical sample, we believe it is essential to perform thorough and robust analyses to be confident with the resulting classification and to suggest tractable strategies to tackle even larger samples. These subsets were obtained simply by arbitrarily splitting the full sample assuming that the order within the SDSS database would provide enough randomisation of the galaxy diversity. We summarise our approach in Fig. 1 as follows:

(i) The full sample with 702 248 spectra was only partially analysed, because the very good agreement between all our analyses did not seem to justify the large computation time that it would have required (Sect. 6).

(ii) Two sub-samples of 300 000 and 302 248 spectra were fully analysed. The latter yields our final classification (Sect. 4), while the former was used for comparison (Sect. 6).

(iii) Seven sub-samples of \( \approx100\,000 \) spectra were used for a first exploration of the data set and a search for the best statistical model (see below). They were also used as a robustness assessment (Sect. 6).

With the 100 000 subsets, the best statistical model is found to be DLM2aju (see Sect. 3.1). Since almost all the other models did not even converge towards any classification, we adopted this statistical model in all the analyses performed for this paper.

We also found that when increasing the number of classes \( K \), the number of very small classes increases while the two to three larger classes remain nearly untouched. As an example, for a sample of 100 000 spectra and \( K = 2 \), the distribution of the number of spectra in the two classes is 94 654, 5346, for \( K = 3 \) we get 86 056, 13 222, 722, and for \( K = 50 \) we get 74 007, 22 129, 1324, 757, 669, 204... with ten classes that only have one spectrum. This behaviour was also found on the larger subsets.

We thus decided to first cluster each subset into three classes, and then apply the Fisher-EM algorithm separately on the two

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**Fig. 1.** Summary of all the clustering analyses performed on different sub-samples. The size of each sample is given in the boxes, and the sizes of the different classes are given in the first two columns to the right, with the percentages given in parentheses. The optimum number of classes is given in the last column to the right. The three topmost samples required a pre-clustering with \( K = 3 \) before the first clustering also with \( K = 3 \) on the largest class gathering more than 99% of the spectra. The analysis of the full sample (702 248 spectra) was not pursued after the second pre-classification for excessive computation time. The four main classes A, B, C and D are indicated in boldface.

(i) The full sample with 702 248 spectra was only partially analysed, because the very good agreement between all our analyses did not seem to justify the large computation time that it would have required (Sect. 6).

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(iii) Seven sub-samples of \( \approx100\,000 \) spectra were used for a first exploration of the data set and a search for the best statistical model (see below). They were also used as a robustness assessment (Sect. 6).
largest classes to obtain a more refined classification. The sub-clustering process can be pursued as much as required by the purpose of the classification and its desired granularity. This procedure is similar to the “canopy” approach proposed in De et al. (2016) in which a first rough clustering is performed before a more refined analysis. Because the Fisher-EM algorithm is much more precise and discriminant than the k-means tool used in De et al. (2016), we often end up with a few classes that have only a very small number of spectra even down to one single spectrum.

This procedure is depicted in Fig. 1 in the column labelled “First clustering” (hence with $K = 3$) while the last column gives the optimum number of clusters obtained by the sub-clustering analyses of the three first clusters. However a supplementary step with $K = 3$, shown by the column labelled “Pre-clustering” in Fig. 1 was necessary for the three samples with 702 248, 302 248 and 102 248, thus containing the same 102 248 spectra. This is explained in Sect. 5.

One can wonder why we need the sub-clustering of classes and why we do not find initially the total number of classes. This is because the clustering made by Fisher-EM is defined in a latent subspace (grossly speaking a subset of wavelengths), which can be different at each stage. There is a specific subspace that discriminates the main classes; as shown later on in this paper, these classes correspond to the gross categories of blue and red galaxies. Then there are different latent subspaces to discriminate between the sub-classes within the main classes.

4. Results

In this section, we present the detailed full analysis of the subset of 302 248 spectra that yield our proposed classification. The other subsets are discussed in Sect. 5.

4.1. Four main classes

A first analysis of the 302 248 spectra with $K = 3$ yields the following distribution: 301 224, 990, 34 (Fig. 1). The smallest group is made of very peculiar spectra that we discard in this study, leaving 302 214 spectra.

The second group (990 spectra) is not sub-clustered any further since its size is not too large and appears to be relatively homogeneous (Fig. 2); it gathers spectra with very blue continua and a lot of emission lines and is called class D in the rest of this paper.

The largest group is then analysed with $K = 3$ leading to the following distribution: 231 604, 64 480, 5140 (77%, 21% and 2% respectively; Fig. 2). The two first classes (that we call A and B, respectively) are different in the slope of their continuum and the emission lines, and the third class C looks more diverse (Fig. 2).

We end up with four main classes (A, B, C, and D). We now present the sub-clustering analysis of first three of them (Fig. 1). The sub-classes are named “$Xn$”, where $X$ is the main class (A, B or C) and $n$ increases as the number of spectra within the sub-class decreases.

4.2. Sub-clustering of class A

The curve of the ICL value (Sect. 3.1) as a function of $K$ (Fig. B.1) shows a strong increase and a significant inflexion at 50 before continuing with a flatter slope. Adopting a larger number of classes does not increase very much the ICL value, and brings many very small classes with less than 10 spectra. The optimum can thus be chosen at $K = 50$. The distribution of the number of spectra among the sub-classes (Fig. 3) decreases nearly linearly in logarithmic scale, hence very sharply: the two most populated classes have 81 992 and 52 044 spectra (35% and 22%, respectively, of 231 604), the other all being below 20 000.

The 50 sub-classes of class A are shown in Fig. B.4. The homogeneity (low dispersion) of the sub-classes is very good for the most populated ones, and tends to degrade for small subclasses. We can already pinpoint very peculiar classes, such as A40 and A50 which probably contain very noisy spectra with some artefacts. Some others seem to contain strange features, such as A4 or A19, which have some strong absorption lines.

4.3. Sub-clustering of class B

The ICL curve peaks at $K = 25$ (Fig. B.2). The distribution of spectra (Fig. 3) shows a dominant class (40 040 spectra, 62% of 64 480), far larger than the others (8592, 4727, 4029, 2785, 1215...) with 5 very small classes of less than 7 members, two having only 1 spectrum.

The 25 sub-classes (Fig. B.5) are nearly all very homogeneous. Three sub-classes (B5, B9 and B11) show a redder continuum reminiscent of the class A, but they have strong emission lines that explain their presence here. The three classes B7, B8 and particularly B13, have wide emission lines.

4.4. Sub-clustering of class C

The ICL curve shows an inflexion at around $K = 10$ (Fig. B.3) that we take as the optimum for the same reason as for class A above. For $K = 15$, 3 classes have less than 4 members, so that we do not consider it a better classification. The distribution of spectra among the sub-classes (Fig. 3) shows a dominant class (3052 spectra, 59% of 5140); the others have less than
700 members. This main class is very homogeneous like two similar but smaller classes (C6 and C7; Fig. B.6). These three classes differ in the detail of the emission and absorption lines, and they also are the only classes with a blue continuum. The other classes are more dispersed, and sometimes peculiar.

4.5. Final classification

The sub-classes of the four main classes A, B, C, together with class D, all based on the 302 248 subset of spectra, builds up our proposed classification in a scheme of 86 classes (Fig. 4). They are organised according to their number of spectra. The largest class, A1, represents 27% of the 302 214 spectra, and the following classes 17% (A2) and 13% (B1), respectively. Class C1 (1%) is at the 14th rank, and class D (0.3%) at the 22nd. The 22 more populated classes are thus sufficient to cover the four main categories that we have identified and encompass 96% of the spectra, while 99% of the spectra are contained in the first 37 classes (up to class B9 which has 319 spectra, Fig. 5).

5. Analyses of the other subsets

For comparison and robustness estimation, we also performed complete analyses of the other set of 300 000 spectra and of the 100 000 subsets. For the full sample of 702 248 spectra, we only performed the pre- and first clustering, both with \( K = 3 \) to limit the computation load (see Sect. 3.3). We summarise the results in Fig. 1.

The previous section shows that the first analysis with \( K = 3 \) on the 302 248 subset yields a very small group of peculiar spectra. It is remarkable that the full sample and that with 102 248 (included in the 302 248 spectra) reveal the same behaviour contrarily to all the other subsets. In other words there are some peculiar spectra contained and individualised in the 102 248 sample that are also individualised in the larger samples that contain them. However, they cannot be the same since there are 7 in the 102 248 subset, and 34 and 493 in the larger subsets. Even though they always represent a tiny fraction of the samples (less than 0.07%), they would deserve further investigation, which we reserve for a subsequent paper.

One lesson of this finding is that there are very few spectra that are different from the vast majority. On one side this is unfortunate since they tend to disturb the clustering analysis imposing a process in several steps somehow depending on the subset. On another side, this demonstrates the discriminative power of the algorithm Fisher-EM which can be explained at least partly by the use of a latent subspace (Sects. 3.1 and 3.3).

For the three sets above (102 248, 302 248 and 702 248), the second classes are considered as being class D as confirmed by the similarity of their spectra. This implies that this class D is absent from all the other subsets.

The analyses with \( K = 3 \) of the largest class of the three sets above and of all the other subsets (of 300 000 and 100 000 spectra) yield very similar distributions of spectra in the three classes that can easily be identified as the main classes A, B, and C.

The optimum number of classes within each of the main class is somewhat different from one subset to the other, but is remarkably consistent. We decided not to further analyse classes of less than about 1500 spectra, which explains why in Fig. 1 the number of class is given as 1 (right column). This concerns class C only. There might be several reasons for the mismatch between the optimum number of classes, but we can point to two main explanations. A first one is because the diversity is not homogeneous between subsets of spectra (they were designed arbitrarily), hence the classes are all not expected to be represented. A second explanation may come from the probabilistic nature of the Fisher-EM classification leading to some fuzziness in the boundaries between classes.

We conclude for a very good consistency between all these analyses and, as shown is the next section, there is a very good agreement between all the obtained classifications. For that reason, investing a large computation time in a thorough analysis of the full sample with 702 248 spectra would not bring much more information than the classification obtained on the subset of 302 248 spectra as presented in Sect. 4.5. The procedure presented in this paper may serve as a guide for other unsupervised clustering of larger samples.

6. Robustness estimation

In this section we assess the robustness of the proposed classification into 86 classes through a five-fold cross-validation technique and using the analyses of the different sub-samples presented in Fig. 1 (Sect. 5).

For this purpose, we first present the two supervised techniques that we used.

6.1. Supervised classification

To assess the robustness of our classification, we need to match the classifications obtained with different samples, that is we
need to classify a spectrum (typically the mean spectrum of a class) into the true classification obtained with the training sample. This is called supervised classification, and we used two techniques.

The first technique uses the statistical model found by Fisher-EM with the training sample to compute the likelihood of the test spectrum on all classes running the E-step of the Fisher-EM algorithm. In other words, we compute the probability that the test spectrum belongs to each of the classes. This process is extremely fast and can be easily applied to classify millions of spectra.

The second technique is the $k$-nearest neighbour (kNN) algorithm. The (Euclidean) distance of the test spectrum with all the spectra of the training sample is computed and the class is determined from the majority vote of the $k$ closest spectra. Obviously this method takes much more time than the previous approach. In addition, the Fisher-EM classification is obtained in a subspace of the features (wavelengths), while kNN uses all features to compute the distance between spectra. However, the kNN algorithm is useful when there is no statistical model for the true classification. If the true classes are available only through their mean spectra, then $k = 1$. When the class attribute is given for the whole training sample, then $k = 5$ is a standard choice and is adopted in this study.
The error rate provided in the R package `mclust` is used in this paper as an indicator of the quality of the match between two classifications. This rate gives the proportion of misclassified spectra in a minimum error mapping (correspondence) between the predicted classification and the true classes. We note that the comparison between our classifications is hampered by the relatively high number of classes (nearly 100) and their extreme imbalance (the classes have from one up to several tens of thousands of spectra).

Hereafter, the reference classification is that obtained with the reference sample of 302,214 spectra and corresponds to the 86 classes shown in Fig. 4. This should not be confused with the true classification that depends on the training sample.

### 6.2. Cross-validation

We performed a five-fold cross-validation in the following way. The reference sample of 302,214 spectra was split into five disjoint sets of about 60,000 spectra. For each such set, we performed the full Fisher-EM analysis on the remaining 240,000 spectra (the training sample) and used the resulting statistical model to predict the classes of the 60,000 spectra (the test sample). We finally compared the predicted classes to the reference classification.

The optimum number of classes for the five cross-validation subsets of 240,000 spectra is between 74 and 116 (as compared to 86 for the reference classification), and the error rate on the five 60,000 sets is between 25% and 42% with an average of 34%. This is called the test error.

We also performed some other tests. For instance, we estimated the training error by predicting the class of each spectrum of the reference sample (302,214) using the statistical model of the reference classification. This error is found to be zero showing the quality of the classification despite the large number of small classes.

We also applied kNN with $k = 5$ on 100 sets of 1000 randomly selected spectra out of 302,214 and found a mean error rate of 11%. We could not do the test on the full sample for excessive computation time.

### 6.3. Comparison with the analyses of other subsets

#### 6.3.1. Distinct sets of 100,000 spectra

The optimum number of classes in the subsets of 100,000 spectra is between 21 and 61 (Fig. 1), hence always lower than 86. This may indicate that the diversity of spectra in the training sample is not represented in all the 100,000 subsets. To ensure a better match between the classifications, we first need to reduce (merge) the reference 86 classes to the number of classes for each subset. This is done with the function `mapClass` from the package `mclust` in R or by matching the mean spectra of the classes with kNN ($k = 1$).

The error rate is found to be between 9% and 21% with a mean at 15% for the six subsets of 100,000 spectra and the subset with 102,248 spectra.

#### 6.3.2. A distinct set of 300,000 spectra

The optimum total number of classes 86 as for the reference classification, but obviously the classes are not exactly the same since the class D is absent (Sect. 5). By matching the number of classes as previously, the error rate is found to be about 30%.

Even though this result is quite satisfactory because the reference and test samples are of similar sizes, it is intriguing that the error is higher than for the smaller samples with 100,000 spectra. We find a clear correlation (Pearson coefficient of about 0.9) between the number of classes and the error rate. Our interpretation is that the culprit is probably the large number of small classes: half of these gather less than 1% of the sample. By construction, the ICL criterion (Sect. 3.1) avoids overfitting as much as possible and favours well-separated clusters (Biernacki et al. 2000), but the nature of very small classes in the context of unsupervised clustering is not so clear. We wonder whether they are due to some kind of over-fitting, are real outliers or are peculiar objects heavily affected by artefacts. Unsupervised classification is a guide towards understanding the sample and only a detailed physical analysis of the spectra in the small classes can answer these questions.

### 6.4. Full sample of 702,248 spectra

Since we did not perform a sub-clustering of the four main classes A, B, C, D, we estimated the error rate between the common set of 302,248 spectra only for these and found 2.2%. This shows the strong significance of the four main classes that are found in all subsets. We infer that the four main classes are four very general categories of the galaxy diversity, as is confirmed by their physical interpretation (Sect. 7).

Since the error rate between our reference classification and the error rates obtained from any subset of the full sample is less than 30%, and given the large size of our subsets and the large imbalance of the classifications, we conclude that the 86 classes obtained from the subset of 302,248 spectra are fully representative of the full sample of 702,248 spectra.

Our strategy to study several subsets of the full sample is successful in estimating the robustness of the proposed classification. This gives us confidence in using this approach for the analysis of other large samples. Instead of simply increasing the sample size to get stuck into computation issues, it is more efficient to study several smaller subsets to identify one that is representative of the full sample. In our case, it seems clear that the subset of 300,000 spectra does not give more information than that with 302,248, which uniquely contains the main class D. The important point is that the representativeness is not given by a priori hypothesis on what the diversity should be, but on purely clustering arguments.

Finally, robustness does not mean that every single class is exactly reproducible. We are working in a very high dimensional space, in which we are looking for the best optimum we can find, not the best in absolute terms if it exists. Smaller classes are probably less robust than the larger classes, but are not necessarily less interesting.

### 7. Discussion

While the optimum number of classes is guided by an objective criterion (here the ICL value), there is still some arbitrariness in deciding how deep sub-clustering must be performed. We have seen that subdividing a class reveals more subtle categories, but it often produces very small classes with a handful of elements. The question arises as to whether this is useful. Naturally, this depends on the purpose of the study. There are fundamentally two goals: performing a supervised classification, that is deciding the nature of a new spectrum, or identifying peculiar objects. The algorithm Fisher-EM is able to tackle the two goals simultaneously by selecting categories during several steps and
focusing on particular categories. In the following, we keep in mind both goals by ensuring that we keep peculiar classes in our study and that we drop out meaningless small classes.

7.1. Description of the resulting classes

We identified 86 classes, but a few of these can be discarded. For instance, the three smallest classes with only one spectrum (A50, B24 and B25) are clearly noisy (see Figs. 4, B.4 and B.5). This also seems to be the case for classes with less than six spectra (A48, A49, B22 and B23). However, some very small classes look rather homogeneous such as B21 with seven spectra. It would certainly be useful to look at individual members of this class. On the other hand, the more populated class A40 can be discarded as well since it clearly gathers data with an artefact around 5000 Å (Fig. B.4).

Apart from these noisy, inhomogeneous, and peculiar classes, most of the classes are very homogeneous (i.e., the mean, 10% and 90% quantile spectra are very similar), but in some cases, the intra-class dispersion is large as it is the case of most of the C sub-classes. This behaviour is especially important in the red part of the spectra, for example, in the case of A31, A32, A34, A36, A41 or B11 (see Figs. B.4 and B.5).

We should note that the dispersion around the mean spectra of the classes (Fig. 4) is always larger than about 10%, hence much higher than the Galactic extinction we did not correct for (about 3% in flux ratio of $g$ to $r$). We cannot exclude some extreme cases that may impact our most dispersed and/or less populated classes which in any case will deserve further investigations in a subsequent study.

7.2. Comparison with other atlases of spectra

In this section, we compare our classification with the atlases by Kennicutt (1992), Dobos et al. (2012) and Wang et al. (2018) described in the introduction (Sect. 1). The spectra of the comparison atlases were resampled to match ours and normalised to the mean between 4300 and 5000 Å. The Kennicutt (1992) spectra are shorter than ours, up to 6983 Å instead of 7371 Å, so that our spectra had to be shortened by 10% in the comparison with this atlas. We must also recall that our spectra are not corrected for galactic dust extinction contrarily to the atlas spectra. But we here use the mean spectrum of each class, averaging out the extinction effect which is nevertheless small as compared to the intra-class dispersion (Sect. 7.1).

Our classes are first matched with these atlases using the kNN ($k = 1$) algorithm in Sect. 7.2.1. The reverse, that is the match of the atlas spectra with our classes, is done with the statistical model provided by Fisher-EM and presented in Sect. 7.2.2.

7.2.1. Match of our classes with the atlases

The matches between each of our 86 classes with those of the atlases with the kNN ($k = 1$) algorithm (Sect. 6.1) are shown in Figs. C.1–C.3. The fit is generally good except for a number of spectra (see below). We must stress that since we have a much larger sample and more classes than these atlases, it is expected that many spectra and classes could not be present in the atlases, hence the fit is sometimes approximate.

As can be noticed in the figures, our spectra are much better fitted by the Dobos et al. (2012) atlas (see Fig. C.2) than the other two atlases (Figs. C.1 and C.3). For instance, the mean of the difference between the fitted spectra is always below 15% (average 5%) with an average standard deviation less than 20% for the Dobos et al. (2012) atlas, except for our class B21 which is matched with a star-forming galaxy spectrum (SFI_0). The agreement with the Kennicutt (1992) atlas appears to be less good. The differences are always lower than 18% (average 5%) except for classes D and B21 because the standard deviations are on average 20%. For the Wang et al. (2018) atlas, the mean difference is lower than 31% (average 7%) and an average standard deviation of 20%.

This ascertainment is foreseeable since the Dobos et al. (2012) atlas is the most representative of the diversity of galaxy spectra comprising 32 classes and spanning three colours (red, blue and green) and six nuclear and star formation activities (from passive to Seyfert galaxies, see their paper for more information). On the contrary, the two other atlases are either very poor in the number of classes or in the number of spectra representing a class. For instance, the Wang et al. (2018) atlas comprises six classes spanning nuclear and star formation activity and the Kennicutt (1992) atlas is made of individual spectra (not averaged spectra) representing a single class. Hence the noise in the spectra of Kennicutt (1992) atlas is higher and can disturb the kNN analysis. Moreover, the individual spectra in this atlas correspond to the classes defined only by the morphology of the galaxies.

In general, when the fit is not of good quality, the mismatch occurs in the red part of the spectrum where our spectra (in black) are often redder than those of the three atlases (shown in red). The disagreement is most important in the fit of subclasses of our class C, which is foreseeable since the dispersion in this class (and sub-classes) is very important; this can be seen in Fig. B.6 where the dispersion is represented by the spectra of the 10% and 90% quantiles. Also the mismatch in general may occur in the emission lines, probably ruling out the extinction correction difference between our spectra and those of the atlases. This behaviour seems to be more important in the fits with spectra from Wang et al. (2018) atlas. This is also foreseeable, since the spectrum representing a class in an atlas (except for the Kennicutt 1992 atlas) is the average of the spectra in that class. As a matter of fact, the line intensities and ratios

![Fig. 6. Distribution of our classes A, B, C and D among the classifications by Kennicutt (1992), Dobos et al. (2012) (see nomenclature explanation in Table C.1) and Wang et al. (2018) from top to bottom.](image-url)

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in averaged spectra are most probably not preserved. In addition, the kNN technique does not take the intra-class variance into account. In the case of the Kennicutt (1992) atlas, a spectrum representing a class is an individual spectrum preserving the physical line ratios and intensities, but it corresponds to a given galaxy morphology that cannot be representative of the wealth of a galaxy class.

The corresponding contingency table (Table C.1) summarises the results of the fits of each of our classes with the three atlases. Our 86 classes are distributed into the 6 classes of Wang et al. (2018), but in only 12 of the 24 classes of Kennicutt (1992), of which the missing classes are cD, cE, dE, IBm, Im, Sb, SBa, SBb, Sbc, SBm and Sm. These are also distributed into 26 of the 38 classes of Dobos et al. (2012), of which the missing classes are GG, h_G, h_GG, h_RG, hh_BG, l_GG, l_RG, p_RG, RED0_0, RED1_0, RG and t_BG (see Table C.1 for the meaning of their nomenclature). As mentioned before, in the Kennicutt (1992) atlas, the categories are defined solely from morphology, not spectral features, while the Dobos et al. (2012) atlas has many overlapping categories.

7.2.2. Match of the atlases with our classes

Conversely, with the use of the statistical model provided by Fisher-EM, we classified the spectra of the three atlases into our own classification. This is shown if Figs. C.4, C.5 and C.6. We also performed a kNN ($k = 5$) fit and the result is quasi-identical. For the Dobos et al. (2012) and Wang et al. (2018) atlases, the
agreement is pretty good, apart from some emission lines that are clearly not matched well. For the Dobos et al. (2012) atlas, the difference in the match has a mean always lower than 4% with a standard deviation of less than 8%, except for the class SF0_0 for which these values are about 9% and 13%, respectively. For the Wang et al. (2018) atlas, these values are 4% and 6%, respectively. Yet, our class A1 seems over-represented.

The match with the Kennicutt (1992) atlas is less good, and is particularly bad for the spectra cG, I, and Sm, which are very noisy, and those of dI, Im, S, S0 (2 spectra), Sb, and SBm, which are all best fitted with our D class or one sub-class of our C class. This result shows that our C class is mainly defined by strong emission lines. As a matter of fact, the spectra of the Kennicutt (1992) atlas that are less well fitted are those with strong emission lines and could only be fitted (badly) with those of class C and D which are characterised by strong emission lines. Apart from these classes, the differences have a mean of less that 6% and a standard deviation of 34% at most.

Figure 6 provides a summary of the distribution of the atlas categories with respect to our four main classes A, B, C and D. The class A is diverse in morphology while class B is mainly composed of Sc type galaxies according to the Kennicutt (1992) atlas. Following the Dobos et al. (2012) atlas, our class B is clearly made of star-forming galaxies (plus AGN+HII), class A
gathers many kinds of galaxies with little star formation, and class C is more difficult to characterise as with the Kennicutt (1992) atlas. Finally, the simple Wang et al. (2018) atlas confirms that class B is made of star-forming galaxies such as class D, class C comprises LINERS, Seyferts and star-forming galaxies, and class A gathers everything except star-forming galaxies.

### 7.3. Physical interpretation of the classes

From the above description of our classes and the comparison with other atlases of spectra, we can make an initial attempt to draw a physical meaning of our classes. A much deeper analysis will be made in a future work to provide a more precise interpretation.

Class A is definitely composed of red galaxies, with a gradual activity of their nucleus or star formation. The passive galaxies are represented for example by classes A8, A10, A12, A18 with different reddening of the continuum, a slight activity is represented for example in classes A1, A2, A6, A9, A31, A32... with a gradual strength of the \( H\alpha \) line and an absence of \( H\beta \). A number of sub-classes of class A are clearly representative of LINER galaxies with gradual values of the line ratios [OIII]/H\(_\beta\) and [NII]/H\(_\alpha\), this is the case for example of classes A3, A4, A5, A7, A11, A13, and A14. There are also transition objects at different quenching stages (A13, A14...). Finally, in class A, we find also sub-classes that represent Seyfert 2 galaxies such as A16, A17, A19, and A23.

Class B is made of bluer galaxies with a higher activity deduced from the stronger and higher number of emission lines. This class is representative of different levels of star-forming galaxies (example classes B1, B2, B3, B4, B6... but also seemingly, of Seyfert 1 type galaxies as suggested by the spectra of classes B7, B8, B13 and B21. Finally some sub-classes show redder continuum (but bluer that class A galaxies) with a higher activity than class A. This is the case of classes B5 and B9.

Classes C and D are characterised by strong and more numerous emission lines with the highest activity being in class D (see Fig. 4). Indeed, emission lines dominate the spectra of class D and are all extremely blue. Consequently, Class D is definitely representative of high activity of star-forming galaxies. Class C is defined exclusively by its set of emission lines that represent a transition class between class B and D in terms of activity. Hence, the spectra of class C span a wide range of colours from very red to very blue galaxies. In Fig. B.6, the sub-classes of class C confirm that this class is definitely characterised by the emission lines. As a matter of fact, the width of \( H\alpha \) line seems to define the sub-classes, for example classes C8 and C9 span wide ranges of continua but present a large \( H\beta \) width. This is also the case for classes C2, C4 and C10 showing a single line width each that is different from one class to the other but a large variation in the continuum within a single class. The fact that our method is able to distinguish classes from the line width of their lines (for instance in our case, the \( H\beta \) line) is a very promising and powerful result. Finally, Class C is probably representative of galaxies with both star formation and nuclear activity, but a more thorough analysis of the spectra is needed to interpret its sub-classes.

### 7.4. An illustration of the diversity of our 86 classes

A detailed analysis of the properties of the classes is kept for a subsequent paper, but we wish to present some hints about the distribution of our classes in two famous scatter plots: a colour-magnitude diagram and the BPT (Baldwin et al. 1981) which is often used to discriminate between star-forming and AGN-dominated galaxies, that is to determine the ionisation origin of the emission lines.

The colour–magnitude diagram (Fig. 7) confirms the analysis above based on the spectra themselves: the A class mainly belongs to the red branch, the B class to the blue branch. There are only very few exceptions to this scheme, such as A7 (blue branch) and may be B11 (blue branch). The C class is more diverse while the D class is obviously very blue and faint. This confirms that the main classes A, B, and D are discriminated by the slope of the spectra, whereas the main class C possess other specific characteristics.

The BPT diagram of [OIII]/H\(_\beta\) versus [NII]/H\(_\alpha\) (Fig. 8) shows that A classes are generally situated around \([\text{OIII}]/H\beta \approx 1\), principally as star-forming or LINERs galaxies. The B classes are mostly on the verge between star-forming and other types, confirming that most of these galaxies are probably subject to intense star formation activity. The C classes kind of avoid the star-forming region (lower left zone) but are rather situated in the Seyfert, AGN of LINERs regions. Class D seems very specific, occupying a tiny region along the top left part of the star-forming limit, similarly to B15 or C7 for instance.

These two diagrams demonstrate that our classification obtained with the unsupervised clustering technique called Fisher-EM retrieves the diversity of the physical properties of galaxies with many details. They can usefully be compared with the classification in ten classes obtained by Chattopadhyay et al. (2019) using a very different set of features (photometry, spectroscopy, morphology, chemical composition, and kinematics) and a very different technique (ICA + k-means). Clearly, working with spectra is more direct and objective since the features do not have to be carefully selected, and yield a much more detailed classification.

### 8. Conclusions

Using a latent subspace clustering algorithm called Fisher-EM, we were able to establish a robust and thorough classification of a large sample of spectra of galaxies from the SDSS DR7. The classification has been obtained with 302 248 spectra, in a hierarchical process of sub-clustering the main classes. Cross-validation with the analyses of several subsets of our full sample (702 248 spectra) demonstrates the robustness of our proposed classification.

The first step of classification with few classes always yields four main categories, grossly corresponding to quiescent, star-forming, Seyferts and LINERs, and galaxies with many emission lines. Subsequent clustering of these main classes yields a total of 86 classes, among which many very small classes are present, sometimes made of peculiar spectra, but also probably revealing pars-differe interesting clusters.

In this paper, we did not attempt a detailed modelling of the mean spectra of the classes to derive the physical parameters that could characterise them. This is the purpose of a subsequent work. Instead, we matched our classification with three atlases in the literature by using either the statistical model provided by the Fisher-EM analysis or the simple kNN algorithm. The generally good outcome of the correspondence shows that an entirely automatic approach to build template spectra of galaxies is achievable and providing a way to perform a supervised classification on millions of spectra.

We more or less reach the same conclusion as Wang et al. (2018) when they compare their classification with that by Dobos et al. (2012) since the detailed match for emission lines
is sometimes disappointing. Classifying complex spectra is certainly difficult, as illustrated for instance in Rahmani et al. (2018) in which a semi-supervised algorithm (self-organizing maps) is used to classify spectra against templates with a very small sample. However, we identify two possible caveats related to the use of templates. Firstly averaging spectra may lead to a loss of some precise and crucial information in the lines, thus we think this point should be addressed specifically on large samples. We intend to pursue the present work in this direction. In particular, we are completing two studies on simulated spectra of galaxies, one with simple single stellar populations (SSPs) and one using the CIGALE code (Boquien et al. 2019). Secondly, we believe it is important to take the intra-class variance into account when comparing with templates or other classifications. This is done automatically when using the statistical model provided by the Fisher-EM algorithm or any other kind of probabilistic clustering technique (e.g., GMM algorithms).

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Appendix A: Noise reduction and binning of original spectra

To preserve the shapes of emission and absorption lines as much as possible, each spectrum was first resampled, increasing its size by a factor of two. We then applied a wavelet decomposition before reducing the noise by applying a threshold, that is by cutting the two shortest components that most probably carry the noise (Donoho & Johnstone 1994; Souza-Feliciano et al. 2018). This step required some manual adjustments to set the best threshold levels by comparing some emission lines before and after noise reduction (Fig. A.1). Lastly, each spectrum was rebinned to reduce its length by a factor of 4, yielding 1437 wavelengths (see Sect. 2).

Appendix B: Sub-clustering

The main classes A, B, and C were further analysed with Fisher-EM to find more refined clustering (Sects. 4.1–4.4). We searched for the optimum number of classes using the ICL criterion for each of them independently (Figs. B.1–B.3). For class B there is a clear peak at $K = 25$. For classes A and B, the ICL curve presents a strong increase at low $K$s and then flattens significantly while continuing to increase slightly (by a few percent). We were not able to reach a peak and the results for larger $K$s yielded more and more very small classes (1–3 members) while the larger classes do not change much. For instance in the case of class C, for $K = 60$ the distribution is 2838, 703, 664, 416, 150, 90, 44, 41, 30, 20, ... 3 (8 classes), 2 (10) and 1 (20), so only 11 classes have more than 10 spectra, while for $K = 10$ this distribution is 3052, 654, 516, 395, 295, 116, 54, 21, 19, 18. Because of this and because the value of ICL does not vary much, we adopted the value at the inflexion of the curves as shown by the red dot (i.e., 50 for A and 10 for C). The resulting classes are shown in Fig. B.4–B.6.
Fig. B.4. Sub-classification of class A with 231,604 spectra.
Fig. B.5. Sub-classification of class B with 64 480 spectra.
Fig. B.6. Sub-classification of class C with 5140 spectra.
Appendix C: Match with other atlases

In this appendix, we present the comparison of our classification with some published atlases of templates as discussed in Sect. 7.2.

Fig. C.1. kNN match of our classes to the atlas by Kennicutt (1992). See Sect. 7.2.
Fig. C.2. Same as Fig. C.1 for the atlas by Dobos et al. (2012).
Fig. C.3. Same as Fig. C.1 for the atlas by Wang et al. (2018). “StarForm” stands for Star Forming, Hα− for Hα-weak and “Comp.” for Composite galaxies.
Table C.1. Correspondence between our classes (Fisher-EM) and those of other atlases (Kennicutt 1992; Dobos et al. 2012; Wang et al. 2018).

| Fisher-EM | Kennicutt1992 | Dobos+2012 | Wang+2018 | Fisher-EM | Kennicutt1992 | Dobos+2012 | Wang+2018 |
|-----------|---------------|------------|-----------|-----------|---------------|------------|-----------|
| A1 E1     | G             | Halpha-weak| A25 E1    | RED3_0    | LINER         |            |           |
| A2 E1     | RED3_0        | LINER      | A26 Sab   | t_G       | Seyfert       |            |           |
| B1 Sc     | hh_G          | Star Forming| B14 Sc    | SF1_0     | Star Forming  |            |           |
| A3 I      | l_G           | LINER      | C6 Sc     | SF0_0     | Star Forming  |            |           |
| A4 Sab    | h_BG          | Composite  | A27 Sb    | RED4_0    | LINER         |            |           |
| A5 I      | t_GG          | Halpha-weak| A28 Sb/Sc | l_BG      | Composite     |            |           |
| A6 I      | RED4_0        | LINER      | B15 Sc    | SF0_0     | Star Forming  |            |           |
| B2 Sb     | SF1_0         | Star Forming| B16 Sc    | SF0_0     | Star Forming  |            |           |
| A7 Sa     | h_BG          | Composite  | B17 I     | SF3_0     | Star Forming  |            |           |
| A8 Sab    | p_G           | Passive    | A29 I     | hh_RG     | Composite     |            |           |
| B3 Sc     | SF1_0         | Star Forming| A30 S0    | p_BG      | Composite     |            |           |
| B4 Sb     | p_BG          | Star Forming| A31 Sb    | RED4_0    | LINER         |            |           |
| A9 Sb     | RED4_0        | LINER      | B18 Sc    | SF0_0     | Star Forming  |            |           |
| C1 Sc     | SF0_0         | Star Forming| A32 Sb    | RED4_0    | LINER         |            |           |
| A10 Sab   | p_GG          | Passive    | C7 Sc     | SF0_0     | Star Forming  |            |           |
| B5 Sc     | hh_GG         | Composite  | A33 Sb/Sc | l_BG      | Composite     |            |           |
| A11 I     | RED4_0        | LINER      | A34 Sb    | RED4_0    | LINER         |            |           |
| A12 E1    | RED2_0        | Passive    | A35 S0:   | s_RG      | Seyfert       |            |           |
| A13 I     | hh_RG         | LINER      | B19 Sc    | SF1_0     | Star Forming  |            |           |
| A14 Sb    | BG            | Composite  | A36 Sb    | t_RG      | LINER         |            |           |
| B6 Sc     | hh_G          | Star Forming| A37 Sb/Sc | BG        | Composite     |            |           |
| D S       | SF0_0         | Star Forming| A38 S0:   | s_RG      | Seyfert       |            |           |
| A15 Sa    | SF4_0         | Composite  | A39 Sb    | s_RG      | LINER         |            |           |
| B7 SB0    | s_BG          | Star Forming| C8 Sb     | RED4_0    | LINER         |            |           |
| A16 SB0   | s_G           | Seyfert    | C9 Sb     | RED4_0    | LINER         |            |           |
| A17 S0:   | s_GG          | Seyfert    | A40 I     | RED4_0    | Halpha-weak   |            |           |
| C2 S0:    | s_BG          | Seyfert    | A41 Sb    | RED4_0    | LINER         |            |           |
| B8 Sc     | SF2_0         | Star Forming| A42 I     | hh_RG     | Seyfert       |            |           |
| A18 Sb/Sc | l_BG          | Composite  | C10 S     | s_BG      | Seyfert       |            |           |
| C3 I      | SF4_0         | Star Forming| A43 I     | hh_RG     | LINER         |            |           |
| C4 I      | t_GG          | Seyfert    | A44 I     | RED4_0    | LINER         |            |           |
| A19 Sab   | t_G           | Composite  | B20 Sc    | SF1_0     | Star Forming  |            |           |
| B9 S0:    | s_BG          | Seyfert    | A45 Sb    | h_BG      | Composite     |            |           |
| A20 Sb    | RED4_0        | LINER      | A46 I     | RED3_0    | Halpha-weak   |            |           |
| C5 Sb     | RED4_0        | LINER      | A47 I     | hh_RG     | Seyfert       |            |           |
| A21 Sb/Sc | h_BG          | Composite  | B21 Sc    | SF1_0     | Star Forming  |            |           |
| B11 I     | SF4_0         | Composite  | B22 Sc    | SF3_0     | Star Forming  |            |           |
| A22 Sb    | RED4_0        | LINER      | A49 S0    | SF4_0     | Composite     |            |           |
| B12 Sc    | SF1_0         | Star Forming| B23 Sc    | SF1_0     | Star Forming  |            |           |
| A23 I     | hh_RG         | Seyfert    | A50 I     | hh_RG     | LINER         |            |           |
| A24 Sb    | t_RG          | LINER      | B24 Sc    | SF0_0     | Star Forming  |            |           |
| B13 BD    | SF2_0         | Star Forming| B25 Sc    | SF0_0     | Star Forming  |            |           |

Notes. Our classes are ordered according to their number of spectra. Dobos et al. (2012) define the nomenclature as: \( p = \text{passive}, l = \text{LINER}, s = \text{Seyfert}, h = \text{H}\alpha, hh = \text{AGN+HII}, t = \text{all}, BG=\text{all blue galaxies (idem for GG (green) and RG (red)).}
Fig. C.4. Match of the atlas by Kennicutt (1992) to our classes using the E-step. See Sect. 7.2. The strong discrepancies in the fits for some classes (dI, Im, SO, Sb:) are due to very strong and thin emission lines in the Kennicutt (1992) spectra. The fits are strongly improved when these spectra are smoothed to reduce the noise.
Fig. C.5. Same as Fig. C.4 with the atlas by Dobos et al. (2012).
Fig. C.6. Same as Fig. C.4 with the atlas by Wang et al. (2018).