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A *diff* procedure for music score files

Computation and visualization of the differences between two music score files

Francesco Foscarin

CÉDRIC lab

CNAM Paris

Paris, France

francesco.foscarin@cnam.fr

Raphaël Fournier-S’niehotta

CÉDRIC lab

CNAM Paris

Paris, France

fournier@cnam.fr

Florent Jacquemard

INRIA and

CÉDRIC lab, CNAM Paris

Paris, France

florent.jacquemard@inria.fr

ABSTRACT

Comparing music score files is an important task for many activities such as collaborative score editing, version control and evaluation of optical music recognition (OMR) or music transcription. Following the Unix *diff* model for text files, we propose an original procedure for computing the differences between two score files, typically in XML format. It performs a comparison of scores at the notation (graphical) level, based on a new intermediate tree representation of the music notation content of a score and a combination of sequence- and tree-edit distances. We also propose a tool to visualize the differences between two scores side-by-side, using the music notation engraving library Verovio, and we employ it to test the procedure on an OMR dataset.

1 INTRODUCTION

Algorithms and tools for comparing text files have been around for more than forty years (Unix *diff* [14],[13]). Their purpose is to identify differences between two text files relatively similar (typically two versions of the same file), at the granularity level of lines with the *Longest Common Subsequence algorithm* (LCS), or at the granularity level of characters, with edit distances. They output, in a normalized format, a list of differences between the two files that corresponds to our intuitive notion of difference. This list is called a *patch* or an *edit script* and, combined with either one of the two files, enables the reconstruction of the other file. Another application is to merge two files containing independent changes into a single file. Those tools are widely applied to text based documents nowadays, for software development, collaborative edition and version control systems.

Similarly to text files, comparing music scores is relevant for several applications. At a global level, it helps to define metrics capable of grasping the *distance* between two scores, for example to evaluate the quality of a transcription process (see [8, 19] for recent proposals). At a detailed level, it is very valuable for musicologist and developers of version control systems to get precise clues on the *locations* of the differences between scores (*e.g.*, between two editions of the same score). One difficulty that immediately arises for defining a *diff* tool for music scores is that, due to the nature/complexity of the music language, a music score contains multiple levels[6, 10] that can be compared.

Most of the literature so far focuses on the *musical content* level, *i.e.* the sequence of events in the score that describe the intended production of sounds, independently from any encoding or rendering concern. The comparison is made with some extended *edit distance* [2, 8, 18–20], or employing tree models [4, 23].



Figure 1: Three staves highlighting the differences between comparison at a music content and a music notation level, in the case of simple monophonic excerpts. The staves (1,2) have the same music content but differ in their music notation. The staves (1,3) differ only in one note (a longer C at the end of the first bar) from one music content point of view, but they are very different in term of music notation.

In the present paper, instead, we focus on the *musical notation* level, *i.e.* how the music is intended to be displayed in the score. In particular we consider 7 elements: note-heads (and rest type), beams, tuplets, ties, dots, note position and accidentals that can be found in scores along with musical content (Figure 1).

The authors of [3], following the work of [17], observe that it does not make sense to apply directly the standard Unix *diff* utility to XML score files. A possible solution is to extract a linear representation of the graphical content [6], but motivating by the hierarchical structure of note beaming and tuplet grouping we chose to follow another approach and compare scores in terms of hierarchical structure, by using a *tree-edit distance* based on tree nodes operations, as proposed by [27] or [7].

In this paper we propose a complete methodology for score comparison at music notation level, articulated in four main contributions: (i) the definition of an original tree-based representation of the music notation content of a bar, (ii) a new edit distance algorithm yielding a list of differences between two scores, (iii) a graphical tool to represent two scores side-by-side with coloured annotations and (iv) a visual evaluation on a OMR dataset.

The tree-based representation of the music notation content of a bar (Section 3.1), acts as a canonical model for disambiguation of the rhythmic content of score formats. For instance, in the case of the XML/MEI format, one score can have many different presentations that lead to the same graphical output, *e.g.*, by using or not a nested structure for beaming, interchanging the order of elements, *etc.* The situation is similar for MusicXML and other formats. Moreover, using an intermediate representation allows to decouple our

approach from the exact file format considered. Anyway for practical reason explained in Section 4 we chose to work mainly with scores in XML/MEI format [24].

The new edit distance algorithm is based on our original tree-structured representation and yield a list of differences between two scores first at the bar level (Section 3.2) and then at a finer granularity level (Section 3.3), similarly to a text diff tool that first finds the different lines and then explores the differences between the words of each line. The distance is the size of the smallest list of modifications required to transform one score into the other, and it reflects visual differences between two scores.

The graphical tool (Section 4) represent two scores side-by-side and shows coloured annotations, highlighting the notation differences between the two scores, in a similar fashion as visual diff tools for text. While it is currently based on our algorithm only, it may be extended to present other metrics and should also be easy to integrate into other programs.

The entire workflow is applied to a dataset of scores produced by OMR and allows to easily spot the mistakes made by the transcription algorithm comparing to the manually annotated version (Figure 2).

Related Work. Our aim is to create for music notation a tool similar to the `diff` program by Douglas McIlroy [15], which compares texts. Besides its practical usage, the latter has motivated theoretical studies which led to efficient algorithms for edit distances computation (e.g., [21, 25]). We draw from this work the process of computing the differences with dynamic programming. Comparing music scores has been the object of previous work with different objectives: evaluation of OMR and automatic music transcription (AMT), collaborative score editing, etc., leading to different approaches to the problem.

Knopke and Byrd [17] pioneered with their "work towards a musicdiff program", focusing on comparing several OMR outputs. They show that a traditional XML `diff` tool cannot be used *as is* on MusicXML, and categorize the comparison difficulties. They also propose a rudimentary visualisation tool. Our workflow is similar to [17] although our objectives differ: we build a tool for users, and put more focus on the second step, with a dedicated representation and tree-edit distance computation procedure.

Recently, the subject has been studied in the MIR context of transcription evaluation. Cogliati [8] present an edit distance similar to the Levenshtein distance, aiming at exhaustive research incorporating the characteristics of music notation into their metric. They set up a very sound evaluation process involving human experts, which highlights how algorithms are biased towards one category of differences or another. Our objectives are different since we are mostly interested in computing and displaying the whole list differences between scores, and not only evaluating an edit distance. McLeod and Steedman [19] improve slightly their work towards the goal of a joint metric for AMT performance, but they still do not take into account typesetting differences.

Aiming at improving collaborative editing of scores, [3] introduce the hierarchic paradigm that we followed, and worked with the Zhang-Shasha tree-edit distance [27]. They bring advances from

theoretical computer science (e.g., [5]) into the music notation community. They present one example on an MEI-encoded file, showing that writing a diff tool for XML music scores should be liable to the problems identified by [17] (i.e. XML coding style differences). We introduce an original tree-based representation to go beyond the problem.

Our last contribution, the graphical tool to visualize differences between scores, is inspired by similar tools for texts. They are now ubiquitous, either being standalone dedicated programs like Meld¹ (cross-platform) or FileMerge/opusdiff (MacOS), or integrated into IDEs (Eclipse, IntelliJ). To the best of our knowledge, our work is the first proposition in the context of music scores.

2 MUSIC CONTENT DIFF

In this first section we consider a simple comparison between two monophonic scores at music content level. Intuitively, its purpose is to be able to compare scores according to *the way they sound*. We use an intermediate score representation similar to a piano-roll, with no overlapping notes.

This approach is not original, but it gives a good baseline to introduce our music notation comparison in Section 3.

2.1 Lossy Linear Score Representation

We assume given an XML score, composed of a single monophonic *part*. From the score we extract a sequence of couples $\langle \textit{pitch}, \textit{duration} \rangle$, where the *pitch* is a MIDI value in $[0, 128]$ (128 added to represent rests) and *duration* is expressed in fractions of beats.

This representation in sequences captures the musical content information of a music score disregarding other music notation elements, such as metric cues indicated by grouping events with beams, ties, dots or chords, and information about pitch spelling. Other representations (e.g. `**kern` [16]) exist with the objective of encoding the musical content, but they also embed elements of music notation. In order to clearly separate the two levels we decided to use a simpler representation.

2.2 String Edit Distance

We compare two parts from two different scores by applying the Levenshtein edit distance to extracted sequences. It is based on the three following edit operations [26] on the above triplets (n represent a triplet as above, and ϵ denotes the empty sequence):

ϵ	\rightarrow	n	insertion of a triplet,
n	\rightarrow	ϵ	deletion of a triplet,
n	\rightarrow	n'	substitution of a triplet by another.

Every such operation $\alpha \rightarrow \alpha'$ is associated a cost value $\delta(\alpha, \alpha')$. We assume that $\delta(\alpha, \alpha) = 0$ for all α , and triangle inequality. The cost of an edition sequence is the sum of the costs of all operations involved in the sequence. The *edit distance* $D(s, s')$ between two sequences of triplets s and s' is the minimal cost of an edition sequence transforming s into s' . It can be computed using the following dynamic programming equations, where $|s|$ denotes the length of s , and $n.s$ represents a sequence made of triple n followed

¹<http://meldmerge.org/>

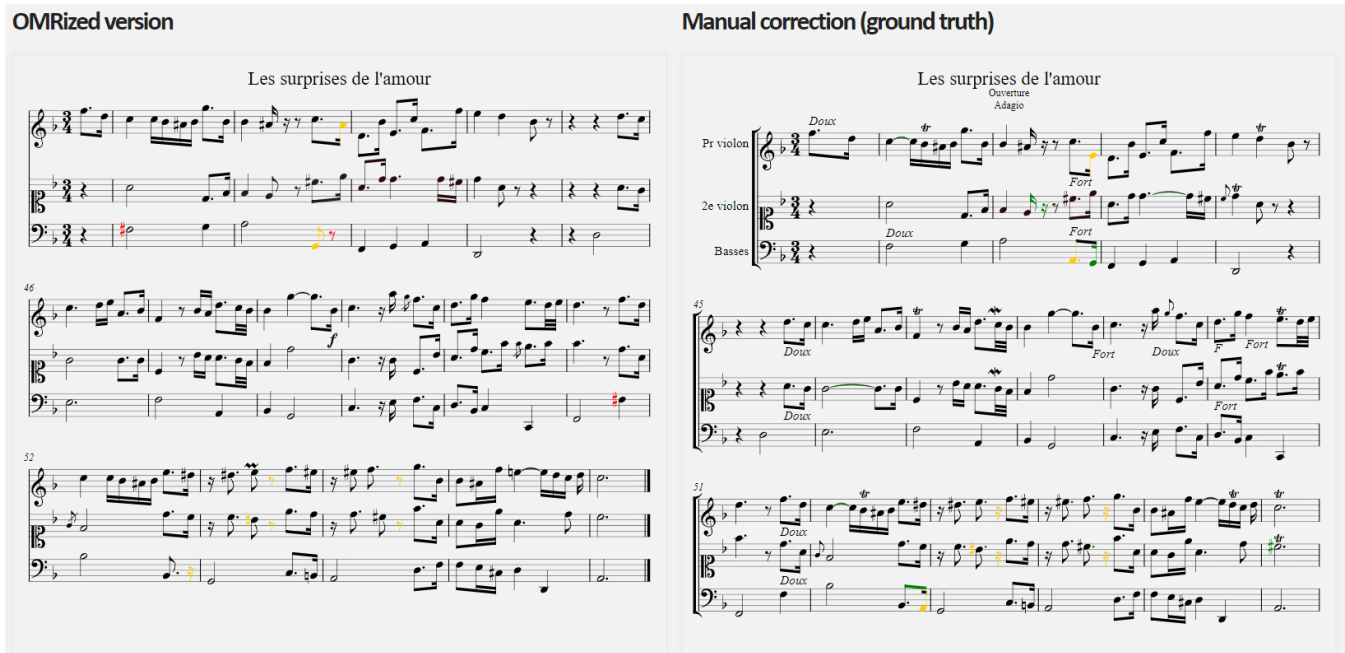


Figure 2: The score diff graphical tool with two scores side-to-side. The differences are annotated with colors: (red) deletions, (yellow) insertion, (green) modifications. In this example it is used to visually evaluate the result of an OMR transcription.

by the subsequence s .

$$\begin{aligned}
 D(\varepsilon, \varepsilon) &= 0 \\
 D(\varepsilon, n'.s') &= D(\varepsilon, s') + \delta(\varepsilon \rightarrow n') \quad (\text{ins}) \\
 D(n.s, \varepsilon) &= D(s, \varepsilon) + \delta(n \rightarrow \varepsilon) \quad (\text{del}) \\
 D(n.s, n'.s') &= \\
 \min \begin{cases} D(s, s') & + \delta(n \rightarrow n') \quad (\text{subst}) \\ D(n.s, s') & + \delta(\varepsilon \rightarrow n') \quad (\text{ins}) \\ D(s, n'.s') & + \delta(n \rightarrow \varepsilon) \quad (\text{del}) \end{cases}
 \end{aligned}$$

3 MUSIC NOTATION DIFF

We now present our main contribution, which is based on another representation, more structured, whose purpose is to enable a comparison of scores according to *the way they look*.

This representation (Section 3.1) is based on tree structures akin to the rhythm-trees for rhythm notation [1, 11, 23], but more tightly related to the music notation information, in order to have an unique representation of the graphical content of the XML scores.

We consider an XML score as a nested structure: on the top level we have a list of score parts (usually one for each instrument), each part is a list of bars, each bar is a list of voices and each voice as a list of *general notes* (notes, rests or chords) that occurs sequentially without overlapping. For the sake of presentation, we will initially focus on scores with a single part, where each bar contains a single voice (Sections 3.1-3.4). We then explain in Section 3.5 how our technique generalizes to polyphonic scores with multiple voices and parts.

The first step is to transform every score into a sequence of trees, one tree (more precisely, one pair of trees, as detailed below) for each bar. Then we proceed in two stages:

- (1) *comparison at the bar level* (Section 3.2), with an alignment of identical bars (*i.e.* they have the same trees), with a Longest-Common-Subsequence (LCS) algorithm.
- (2) *comparison inside-the-bar* (Sections 3.3, 3.4): a more in-depth processing of unaligned bars, using purposely designed tree-edit distances algorithms that descends into the bar representations in order to identify differences.

3.1 Tree-Based Representation

We present an abstract model of the music notation content of bars in music scores. It is designed as an intermediate structure for our algorithms of *diff computation*, and it is not meant to be a format for data exchange. It can be imported from and exported to XML (see Section 4). This model is based on the following two types of tree representations, corresponding to two aspects of music notation.

3.1.1 Beaming Trees. A *beaming tree* represents the notes in a bar and the beams and ties between them. In particular, we encode in the leaves the information about the note-heads, dots and ties; the remaining information (*i.e.* beams and flags) are encoded in the tree structure (Figure 3).

Every leaf of a beaming tree represent an *event*: a note, a chord or a rest. In the XML representation, a chord can be considered in a single voice, if all the notes of the chord have the same duration. Each leaf is labeled by a triple containing the following information (see Figure 3 for an illustration):

- *pitch*s: a sequence where each element contains a *pitch position* (letter + octave number for notes or R for rests), an *alteration value* among 'none' and $\flat, b, \natural, \sharp, \times$ and a *tie flag* indicating if the corresponding note is tied to an element of

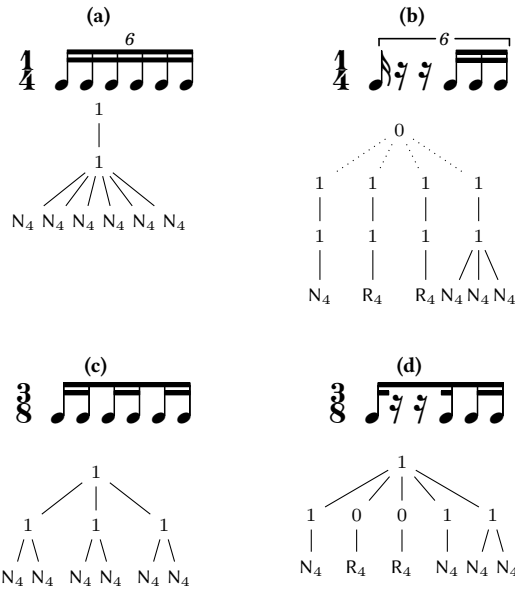


Figure 3: Examples of beaming trees. To simplify the presentation we use a pitch-less notation with notes and rests denoted resp. by N , R . Rests with 2 flags are positioned in the tree like notes with 2 flags, while notes beamed together are represented by a different tree structure.

the previous event in the tree (following a depth first traversal). The pitch position represent the vertical position on the score, not the sounding pitch name, in order for it to be independent from the clefs and transposing instruments; we set it as the pitch name with a G-clef by implementation choice. The sequence is ordered from the lowest to the highest pitch; in case of a single note or a rest (not a chord), *pitches* is a singleton sequence;

- *note-head*: a value representing a fraction of whole note, e.g., 1, 2, 4, represent respectively \circ , \flat , and \blacktriangle for a note, and — , — , z for a rest;
- *dots*: a natural number in $[0, 3]$ that specify the number of dots.

We define the size $\|pitches\|$ of a sequence *pitches* as the sum of the sizes of all its elements, where the size $\|pitch\|$ of an element *pitch* is the number of graphical items it contains, e.g., a note $D4 \sharp$ with a tie will have size of 3, while a rest R will have size of 1.

In the internal nodes, we use the two symbols '0' and '1' for the encoding of *beams*; intuitively a label '1' indicates that there is a beam between the notes in the subtrees under the node. More precisely, let the *weight* of a node n be the number of nodes labelled '1' on the path from n to the root of the tree (including n itself). Let n and n' be events in two successive leaves and ℓ be their least-common-ancestor in the tree (it is always defined and unique). The number of beams between n and n' is either $weight(\ell)$, if in the two paths $\ell \rightarrow n$ and $\ell \rightarrow n'$ all inner nodes are labeled by '1', or it is 0 otherwise.

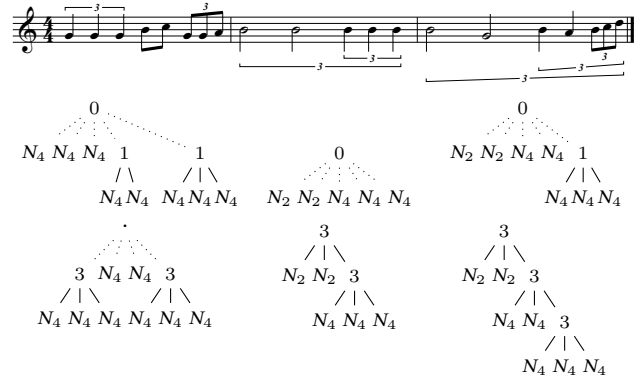


Figure 4: Beaming and tuplet tree for nested triplets (in pitch-less representation). In the first bar, the sequence of the first three notes is labeled by 3 (with a bracket), the sequence of the last three notes is also labeled by 3 (no bracket needed because these three notes are beamed, according to the beaming tree), and the two middle notes do not have a tuplet label.

3.1.2 Tuplet Trees. The *tuplet trees*, illustrated on Figure 4, represent the locations of the numbers and brackets explicitly defining in a score the nature of the triplets. Every inner node of a tuplet tree is either unlabeled (this is denoted with a label \cdot in figures), or labeled by a pair of integers $i : j$, with $i, j \geq 2$, meaning ' i elements in the time of j ', or by a single integer i if j can be omitted [12]. The leaves are labeled like for the beaming trees: Every tuplet tree is associated to a companion beaming tree with the same sequence of leaves (the difference between them is only in the structure).

3.2 Longest Common Subsequences of Parts

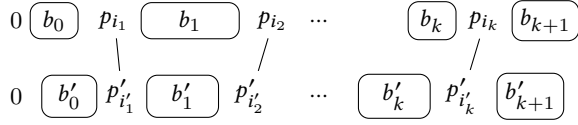
Assume that we are given two parts extracted from two different scores. We represent each part by a sequence of pairs $p = \langle bt, tt \rangle$, one pair per bar, where *bt* is a beaming tree and *tt* is a companion tuplet tree.

Similarly to the line-by-line comparison procedures for text files [14], and considering a bar as the analogue of a line in a text file, we shall align identical bars from the two parts, by computing their longest common subsequence (LCS), whose size is defined by the following recursive Dynamic Programming equations (with similar notations as in Section 2.2):

$$\begin{aligned} LCS(\varepsilon, s') &= LCS(s, \varepsilon) = 0 \\ LCS(p.s, p'.s') &= 1 + LCS(s, s') \quad \text{if } p \equiv p' \\ &= \max(LCS(p.s, s'), LCS(s, p'.s')) \quad \text{otherwise.} \end{aligned}$$

In the second line, $p \equiv p'$ means that the trees in pairs $p = \langle bt, tt \rangle$ and $p' = \langle bt', tt' \rangle$ are identical, i.e. that $bt = bt'$ and $tt = tt'$. According to the representation presented in Section 3.1, it means the corresponding bars can be considered identical.

The above equation can be adapted in order to compute a maximal *alignment* of identical pairs between the two sequences. Formally, denoting the two sequences by $s = p_1, \dots, p_m$ and $s' =$


Figure 5: LCS and difference blocks.

p'_1, \dots, p'_n , the alignment is made of two strictly increasing sequences of indices of same length, $(i_0 = 0, i_1, \dots, i_k)$ and $(i'_0 = 0, i'_1, \dots, i'_k)$ such that for all $0 \leq j < k$,

- (i) $p_{i_{j+1}} \equiv p'_{i'_{j+1}}$, and
- (ii) the pairs in the respective sub-sequences $b_j = p_{i_{j+1}}, \dots, p_{i_{j+1}-1}$ and $b'_j = p'_{i'_{j+1}}, \dots, p'_{i'_{j+1}-1}$ are pairwise distinct (each of these sequences might be empty if $i_{j+1} = i_j + 1$ or $i'_{j+1} = i'_j + 1$).

We call each pair $\langle b_j, b'_j \rangle$ as above a *difference block* (see Figure 5) and the objective of the next two subsections is to perform a fine comparison between blocks.

3.3 Inside-the-bars Comparison

In the following, for a tree t (either a beaming tree or tuplet tree), $\|t\|$ denotes the sum of the sizes of labels of all the nodes n in t (size of labels is 1 by default, $\|n\|$ for a leaf as defined in Section 3.1.1, 0 for unlabeled nodes). For a sequence $s = t_1, \dots, t_k$ of trees, $\|s\| = \sum_{i=1}^k \|t_i\|$.

3.3.1 Comparison of Beaming Trees. The following recursive equations define an edit distance (denoted by B) between sequences of beaming trees. The distance between two trees is defined by the particular case of singleton sequences. In these equations, $t.s$ and $t'.s'$ denote two beaming tree sequences, with $t = a(s_0)$ and $t' = a'(s'_0)$, where s_0 and s'_0 might be empty.

We call a tree *atomic* if it is composed of a single leaf node (i.e. it represents an event). We call a tree $t = a(s_0)$ *unary* if s_0 is a singleton sequence (i.e. the root has a single child).

$$\begin{aligned}
B(\varepsilon, s') &= \|s'\| & B(s, \varepsilon) &= \|s\| \\
B(t.s, t'.s') &= \\
\min \left\{ \begin{array}{ll}
B(s, t'.s') & + \|t\| \quad (\text{del-tree}) \\
B(t.s, s') & + \|t'\| \quad (\text{ins-tree}) \\
B(s_0.s, t'.s') & + 1 \quad (\text{del-node}) \\
& \text{if } t \text{ non-atomic, } t' \text{ atomic, or } t \text{ unary, } t' \text{ not unary} \\
B(t.s, s'_0.s') & + 1 \quad (\text{ins-node}) \\
& \text{if } t \text{ atomic, } t' \text{ non-atomic, or } t \text{ not unary, } t' \text{ unary} \\
B(s_0.s, s'_0.s') & + \delta(a, a') \quad (\text{descend}) \\
& \text{if } t, t' \text{ non-atomic} \\
B(s, s') & + A(t, t') \quad (\text{leaf}) \\
& \text{if } t, t' \text{ atomic}
\end{array} \right.
\end{aligned}$$

where $\delta(a, a')$ is a defined by $\delta(a, a') = 0$ if $a = a'$, and $\delta(a, a') = 1$ otherwise.

The cases (del-tree) and (ins-tree) correspond to the deletion or insert of a whole subtree. The cases (del-node) and (ins-node) with one unary tree correspond to a difference of one beam between the trees. The other cases descend down to leaves and then compare atomic trees t and t' with the distance $A(t, t')$ defined as follows

(p, n, d stand respectively for the attributes *pitches*, *note – heads*, *dots* of Section 3.1):

$$A(t, t') = D(p(t), p(t')) + \delta(n(t), n(t')) + |d(t) - d(t')|$$

where the first literal $D(pitches(t), pitches(t'))$ is the Levenshtein distance defined in Section 2.2, using the following operations costs:

$$\begin{aligned}
\delta(\varepsilon \rightarrow pitch) &= \delta(pitch \rightarrow \varepsilon) = \|pitch\| \\
\delta(pitch \rightarrow pitch') &= \delta(n, n') + \delta(alt, alt') + \delta(tie, tie')
\end{aligned}$$

where n, n', alt, alt' and tie, tie' are the respective name, alteration and tie values of *pitch* and *pitch'*.

3.3.2 Comparison of Tuplet Trees. Tuplet trees have a simpler structure than beaming trees (e.g., no unary nodes). We compare them with the classical Zhang-Sasha equations [27] for computing a tree edit distance (denoted by T), i.e. the smallest number of node-edit operations to transform a tuplet tree into another (instead of the algorithm of Section 3.3.1 which is specific to beaming trees).

$$\begin{aligned}
T(\varepsilon, s') &= \|s'\| & T(s, \varepsilon) &= \|s\| \\
T(t.s, t'.s') &= \\
\min \left\{ \begin{array}{ll}
T(s_0.s, t'.s') & + 1 \quad (\text{del-node}) \\
T(t.s, s'_0.s') & + 1 \quad (\text{ins-node}) \\
T(s, s') + T(s_0, s'_0) & + \delta(a, a') \quad (\text{subst-node})
\end{array} \right.
\end{aligned}$$

where $\delta(a, a')$ is as in Section 3.3.1 for inner nodes and $\delta(a, a') = A(a, a')$ when a and a' are leaf labels.

3.4 Comparison of Difference Blocks

We compute the distance between two difference blocks b, b' using the following recursive equations, similar to the equations of Section 2.2.

We recall that a block is a sequence of pairs (bar representations) of the form $p = \langle bt, tt \rangle$, where bt and tt are respectively a beaming tree and a tuplet tree.

$$\begin{aligned}
D(\varepsilon, b') &= \|b'\| & D(b, \varepsilon) &= \|b\| \\
D(p.b, p'.b') &= \\
\min \left\{ \begin{array}{ll}
D(b, b') & + \Delta(p, p') \quad (\text{edit-bar}) \\
D(p.b, b') & + \|p'\| \quad (\text{ins-bar}) \\
D(b, p'.b') & + \|p\| \quad (\text{del-bar})
\end{array} \right.
\end{aligned}$$

where, for $p = \langle bt, tt \rangle$ and $p' = \langle bt', tt' \rangle$:

$$\Delta(p, p') = B(bt, bt') + T(tt, tt') - \text{corr}(p, p') \quad (1)$$

is the edit distance between two bar representations; $\text{corr}(p, p')$ is a correction of $T(tt, tt')$ to avoid to count the same differences twice (see Section 4).

Finally, the distance between two parts is the sum of distances between all the difference blocks.

3.5 Multiple Voices and Parts

Recall that we consider a model of scores made of a list of *parts* (usually one part for each instrument), where each part is a list of *bars*, each bar is a list of *voices*, and each voice is a list of *general notes* (notes, rests or chords).

The algorithms described in the previous sections deal with the comparison of single-part and single-voice scores. The generalization to polyphonic scores with multiple-voices and multiple-parts is implemented as follows.

multiple-voices: when comparing two bars we consider all couples made of one voice of the first bar and one voice of the second one, we run the algorithm of Sections 3.3 and 3.4 on each couple independently and then take the couples that yield with minimal distance. If the number of voices in the two bars is different, we return the corresponding voice-insertion and voice-deletion.

multiple-parts: we first map the parts in the first score into parts of the second one using the metadata like part or instrument names. We run then our algorithms on each couple of mapped parts independently. In case of parts of score 1 not present in score 2 or conversely, we return the eventual part-insertion and part-deletion. An alternative would be to apply a procedure similar to the above one for multiple voices (coupling parts and taking the ones with smallest distance) but it would be very costly. Indeed, the number of parts can be high (e.g., in the case of orchestral scores), compared to the number of voices in a bar (typically 2 or 3 voices). Moreover, comparing 2 parts require to compare all the data inside these parts (bars, voices *etc.*).

4 IMPLEMENTATION & EXPERIMENTS

The construction of the models of Sections 2.1 and 3.1 from MEI scores, as well as the implementation of the algorithms proposed, have been performed in Python3, on the top of the Music21 toolkit [9].

The computations of the string edit distances of Sections 2.2, 3.2 (LCS) and 3.4 are performed with an iterative computation and run in time $O(m*n)$, where m, n is the length of the two sequences. The computation of the tree-edit distances of Section 3.3, is implemented by recursive functions, with a worse case time complexity in $O(n^4)$. We use a table of solutions in order to avoid to recompute the function on the same inputs and hence reduce the computation time. Nevertheless, efficiency is not a crucial issue in this case, because the difference blocks are typically small (e.g., in OMR evaluation), or such that $b = \epsilon$ or $b' = \epsilon$ (insertion or deletion of bars).

A difference in our score is modeled as a triple (*operation*, t , t') where *operation* is a string identifying the performed operation (e.g. ins-node), and t, t' are the subtrees transformed by that operation (or ϵ is case of insertion/deletion). In the implementation, the above recursive equations are actually extended in order to compute, beside the edit distance, a list of differences. It means that the functions B, T etc. return pairs made of a cost value along with a diff list. To that respect, the operator $+$ in the recursive equations makes the sum of the cost values and the concatenation of *diff lists*, and the min selects the pair with the smallest cost value. The function $corr(p, p')$ in Section 3.4 uses the difference lists returned by B and T , in order to detect intersection that may occur, in particular for leaves. These intersections are removed (operator $-$ in (1)) both from the cost value and the diff list. Thus, we avoid accounting twice for the same difference (e.g., a pitch modification must be counted only once, even if it is present in the result of both B and T).

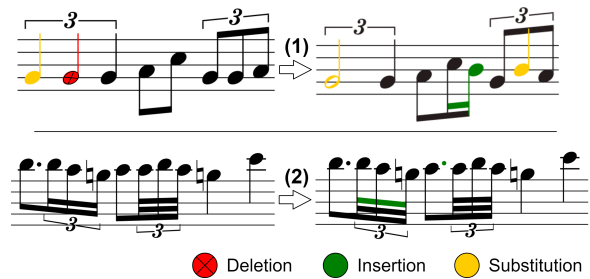


Figure 6: Visualization of differences between two bars computed by the algorithms of Section 3. The upper example (1) is taken from Figure 4 and manually modified to create some differences, while the lower example is a real-life example from an incorrect OMR score import, compared to the correct score.

The elements of *diff lists* are pairs of references to elements in the two scores, plus an identifier of the edit operation. To have an unique reference for the elements in the scores, we use the unique id associated to each element in the MEI format of music scores. We store those ids in the leaves of our tree representation, and they are very useful later in order to display the differences between scores. The XML ids are an important feature, not present in the MusicXML format, and the main reason we always work with MEI score files in input (a MusicXML score can be transformed into a MEI score as a preprocessing).

After computing the edit distances and list of differences between two given scores, we display, using Verovio [22], the two scores side-by-side, highlighting the differences.

We run experiments in order to test the usability of our workflow for two scenarios: collaborative edition of scores, and OMR evaluation. For the first use case, we considered made-up scores with some typical modifications (bar insertions, note pitch modifications, *etc.*) in a context of versioning. For the second scenario, we used a corpus produced by the Bibliothèque nationale de France (BNF) composed of 21 *ouvertures* of Jean-Philippe Rameau, each with an *OMRized* version (from manuscripts using PhotoScore²) and its manual correction, that we compare with our tool³.

Our algorithms has shown promising results in both cases, highlighting very fine differences in the scores (e.g., beams and dots), as illustrated on Figure 6. The evaluation of OMR results were not possible for many scores, due to quality issues in the XML files [10], e.g., a tie between two non-consecutive notes, or syntax errors in the XML file, such as an opened beam not closed. It should be noted that our tree-based model can only handle correctly encoded scores, and that such *faulty* notations will result in an error.

5 CONCLUSION

We have presented a procedures to compare XML scores at music notation level that is based on a tree intermediate representation

²<https://www.neuratron.com/photoscore.htm>

³see our supplementary material page at <https://anonymoususer12042019.github.io/> for details.

of bars and on a combination of string edit distance and tree edit distance.

In contrast with most approaches in the state of the art, our methodology produces a list of differences that can be highlighted directly on the scores, and alleviates the user from the tedious task of manual scores comparison. Our algorithms gives also a similarity metrics, but this metric is only used to compute the list of differences and it is not useful per se. For this reason, comparison to other approaches (e.g., techniques for melodic similarity) would not be relevant.

Experiments show that our approach allows to correctly displays very fine differences in the scores. Technical improvements are still needed in other stages (XML parsing, error detection and correction, complex score visualization) to improve usability.

We leave for future work other graphical aspects that are not considered in the actual implementation, such as stem directions, clefs, slurs, expression markings, ornaments and lyrics. We also plan to investigate on a more complete treatment of multi instruments polyphonic scores, that will correctly handle staves swapping and multi-staves voices.

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