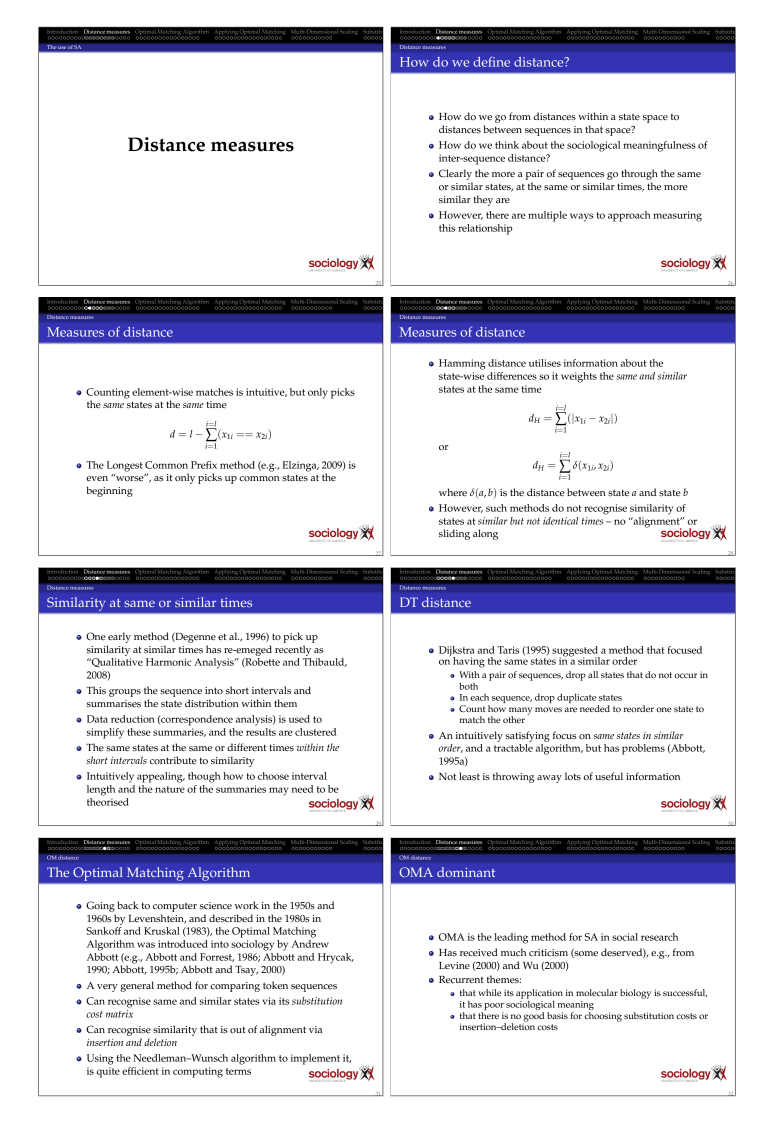
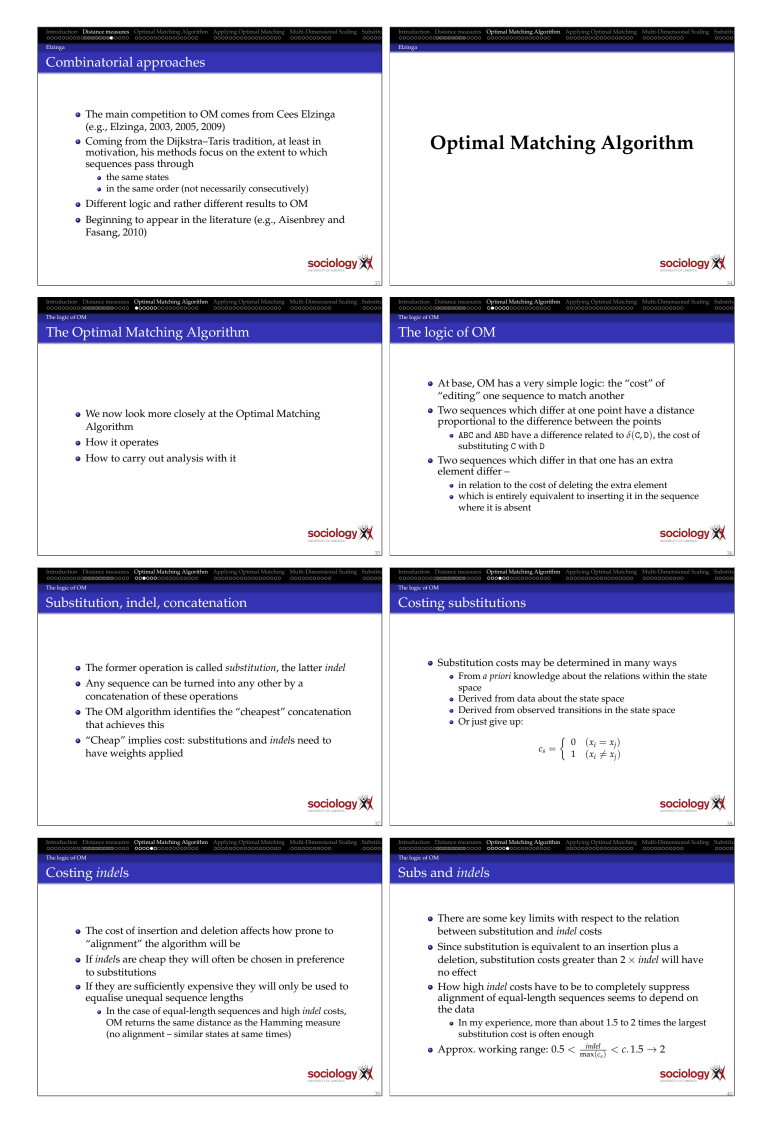
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| Sequence Analysis in Sociology Brendan Halpin Department of Sociology University of Limerick brendan.halpin@ul.ie Helsinki, May 19, 2010 | Introduction |
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| What this workshop is intended to cover | What this workshop is intended to cover |
| Explore the use of Sequence Analysis (SA) in the social sciences, particularly Optimal Matching Explore the analytical use of "empirical typologies" and other measures derived from SA Consider alternatives to Optimal Matching (OM) Provide enough practical information for participants to conduct their own analyses using software How to think about using SA | Explore the analytical use of "empirical typologies" and other measures derived from SA use of OM to generate classifications for further analysis or to generate other sorts of trajectory level information |
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| ^{Overview} What this workshop is intended to cover | Overview What this workshop is intended to cover |
| Consider alternatives to Optimal Matching (OM) Non-aligning methods Combinatorial methods Duration-sensitive methods | Provide enough practical information for participants to conduct their own analyses using software Using SQ add-on for Stata Using my faster but less user-friendly add-on for Stata Using TraMineR for the R statistical language |
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| _{Overview} How to think about using SA | overview Key questions |
| • What's it good for? | Is sequence analysis useful? Does it go beyond exploratory and descriptive? If not, is that enough? |
| When to use other methods?What method to choose?How to adapt it for your problem | What does it "mean"? How do the results inform us about sociological issues? How can we manipulate the inputs to get better meaning? How should we choose between algorithms for different substantive problems? |

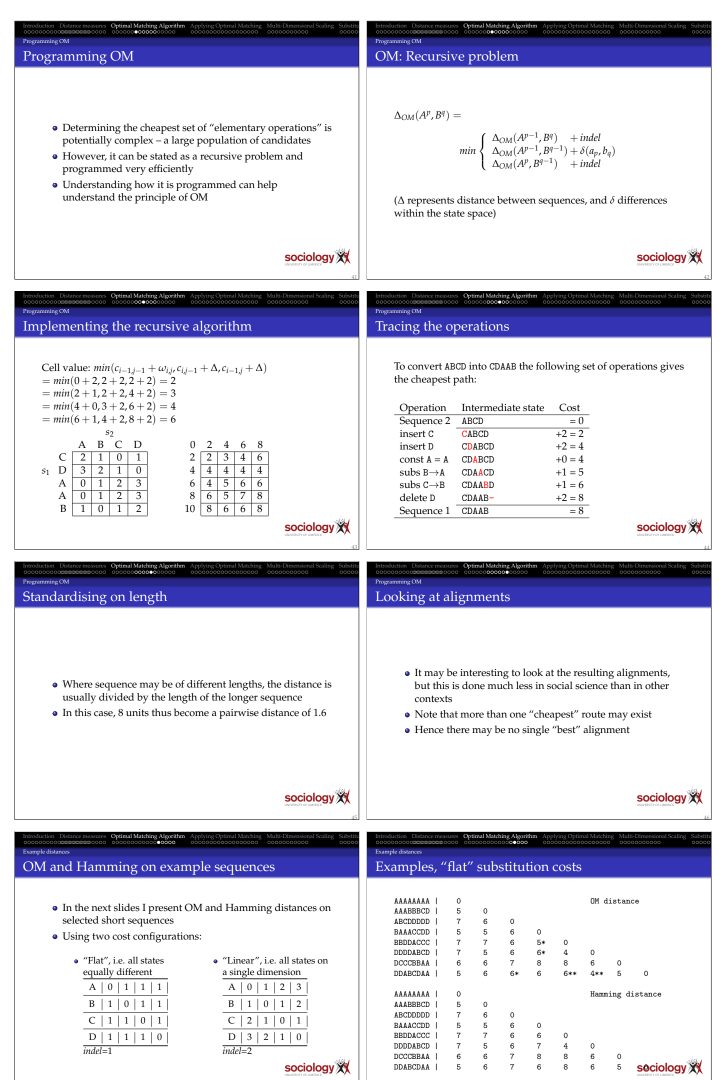
What is Sequence Analysis? What are sequences? • "Sequences" are temporal (or at least linear) trajectories through a state space • Sequences are ordered "trajectories" through a state space • Sequence Analysis treats sequences "holistically" • Their nature depends on • Alternative methodologies are more analytical (and often • the nature of the state space: multi-dimensional, stochastic) and focus on factors such as the generative continuous, real, categorical? processes the nature of the time dimension: atomic, discrete, • Usually at the cost of ignoring some aspect of the continuous, stretchable or rigid? information encoded in the sequence how they start and finish, what they mean substantively In contrast, SA tends to be blind to the processes generating the sequence, thus focusing on the epiphenomenal? sociology 💥 sociology 💥 What are sequences? - the nature of time What are sequences? - state spaces The nature of time has a large bearing on the adequacy of sequences as a representation of the phenomenon "Atomic" sequences consist of elements that are naturally • The state space is important for how we think about distances between points separate and sequential, such as a series of purchases or if multi-dimensional in Rⁿ, point distances are naturally Euclidean or other function of the space votes, or steps or utterances, or CAGT bases in DNA -"time" is naturally discrete Continuous time can be discretised if categorical, we often define pairwise point distances a • If the state changes very frequently (especially if continuous state) we can consider this as "sampling", for example, digitisation of an audio stream priori or empirically (how?) • if many categories? • If change is relatively rare, we may wish to represent the sequence as a series of spells (start and end-times of a period in which the state is constant) sociology 💥 sociology 💥 What is a whole sequence? "Conventional" methods for longitudinal data Many conventional approaches • Hazard rate modelling (event history analysis) • How we define a "whole" sequence is also an important Time-series analysis Loglinear or Markov modelling of transition rates issue - where does it start and end? Start-end tables • For some sequences, it is natural: Panel analysis (cross-sectional time-series analysis, • The steps of a dance, the words of a song multiple-response approaches) The rhetorical structure of a journal article or a folk tale Use of simple summaries of trajectory to predict future • For some purposes, fragmentary sequences can be used outcomes (e.g., searching for DNA matches) • All have analytical strengths - allow inference with respect to clear hypotheses In what way does SA offer something more than they do? sociology 💥 sociology 💥 SA versus conventional Special considerations for life-course sequences • Clearly a classification based on SA will do better than one based on summaries such as start/finish state or • For life-course and other sequences, the requirements of cumuluated duration in states - respects order the analysis impose structure • Relative to hazard rate modelling, SA respects the whole • Usually cannot just match random segments of trajectory, rather than looking at time to a single event employment history (note the existence of repeated events hazard models) We impose comparability criteria (e.g., t_0 is a specific event, follow until a particular outcome or for a specified • Models based on transition rates have difficulty with duration) transition matrices which change in complex ways through Issues of left- and right-censoring become relevant time (due to life course and period effects, for instance) • The various SA methods tend to be blind to these issues Multi-dimensional sequences are even more complex None of the conventional methods offer a digestible descriptive overview sociology 💥 sociology 💥

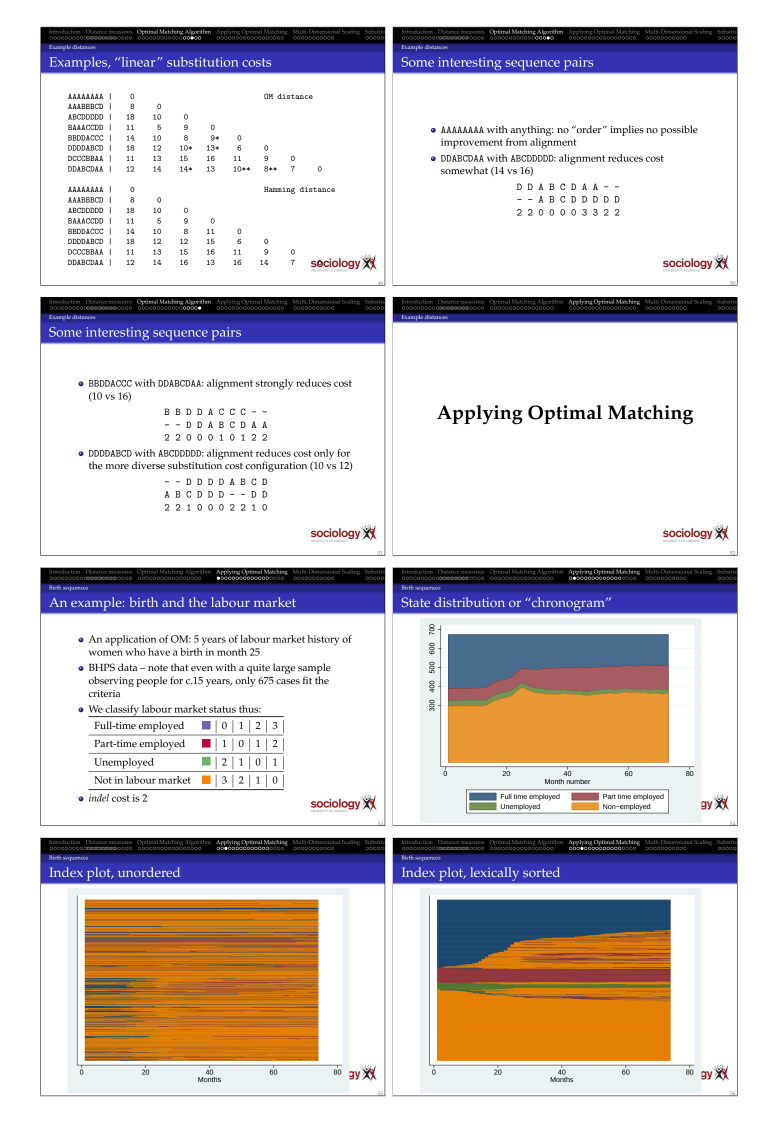
Time consequences Space consequences • For life-course and other sequences, the requirements of • Time that is naturally discrete fits a token-sequence the analysis impose structure representation naturally • Depending on the nature of the state space and the time • "Sampling" continuous time raises issues of distortion due dimension different approaches will be required to the frequency of sampling • A \mathbb{R}^n state space simplifies state distance issues compared • Whether time has a "ruler" or calendar, or is with categorical states, where we need to find a "developmental" or stretchable, has a bearing on how justification for our distances attractive alignment is sociology 💥 sociology XX What do we do with SA: distances (2/2)What do we do with SA: distances (1/2)• SA is very simple: turn information about the state space into information about the trajectories - pairwise distances or similarities • Well chosen ideal-typical sequences can make for very Permits the use of cluster analysis (CA) to generate a interpretable results data-driven classification CA may or may not generate a useful classification Permits comparing all sequences to a set of "typical" sequences • MDS can inform CA, may yield interpretable dimensions itself • Permits analysis of the multi-dimensional space implied by the inter-trajectory distances (multi-dimensional scaling, MDS) • Also permits comparing grouped or paired sequences (e.g., couple's work histories, Han and Moen, 1999) sociology 💥 sociology 💥 What do we do with SA: empirical typology Empirical/theoretical typology • Why is data-driven classification - an empirical typology attractive? • If we can generate a typology of sequences from theory, we • Up to $\sum_{i=1}^{m} n^{i}$ possible sequences, with *n* states and spells may not need SA up to *m* tokens long (e.g., for 4 states over 20 months, more However, can be difficult to write foolproof rules to assign than a trillion possibilities) sequences to groups Observed sequences represent a highly structured subset • SA linking the observed to the ideal typical sequences will • The structure is much more than, say, that summarised by allow us to populate a theoretical typology automatically starting state distribution and Classification by inspection may be possible, but can be the transition matrix averaged over the data set (or even impractical changing through time) • A good classification should (?) pick up this structure sociology 💥 sociology 💥 Empirical typology by CA Exploratory/descriptive: enough? • Cluster analysis on the pairwise distance matrix creates the classification automatically for us · Key questions: can it be more than exploratory and But can be difficult to characterise the resulting groups descriptive? cleanly Is exploration/description enough justification? • And clustering may be unstable if there are not natural groups sociology sociology 💥

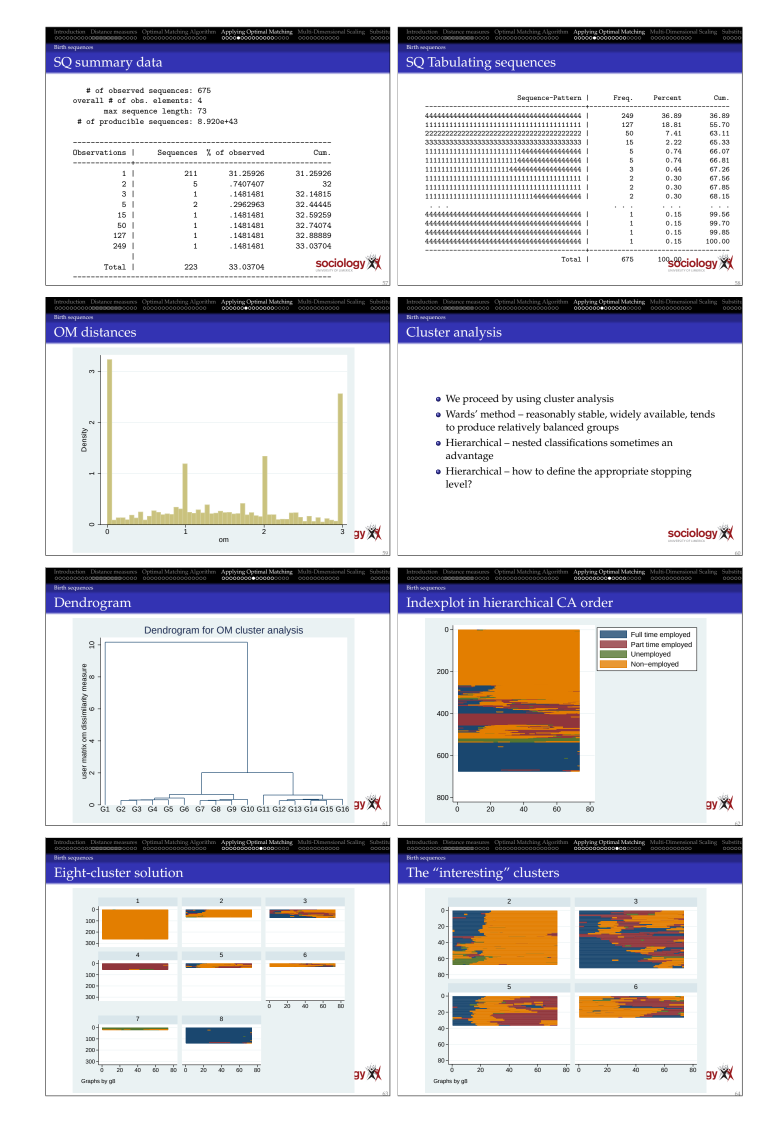
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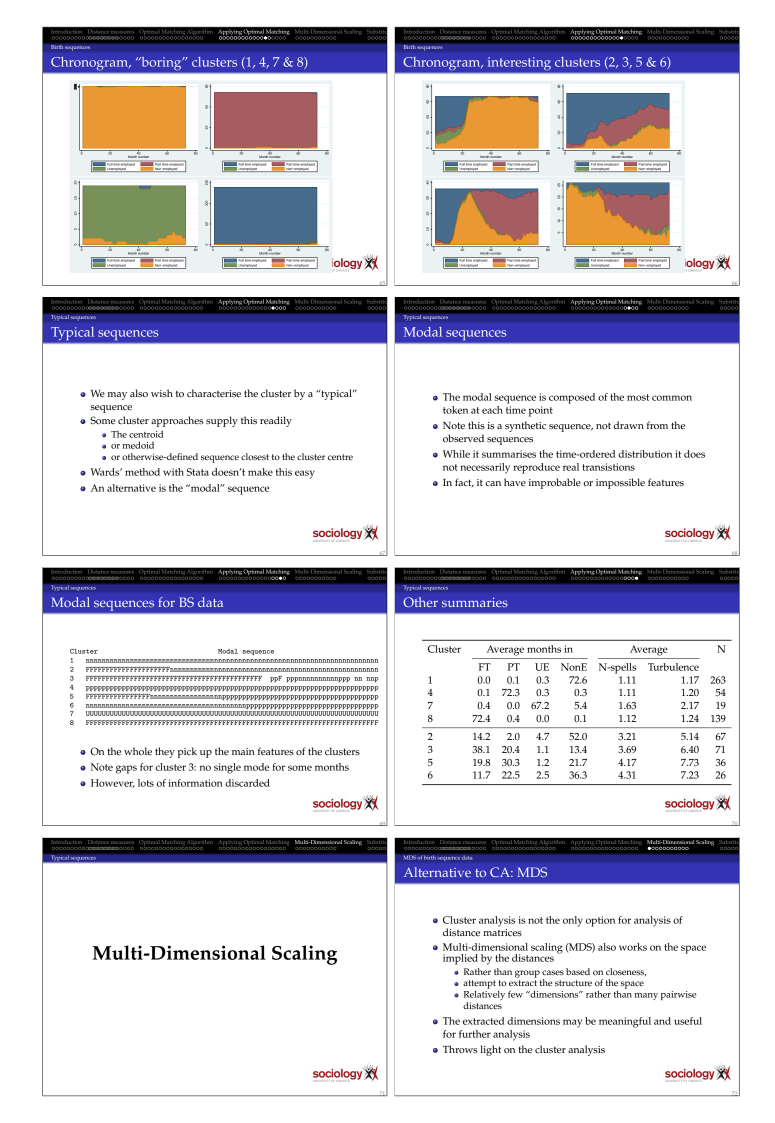










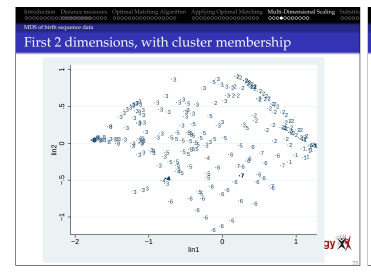


MDS in Stata

MDS of birth

Classical metric multidimensional scaling dissimilarity matrix: omlin

| | | | Number of | | 6 |
|---------------|------------|----------|------------|-------------|---------|
| Eigenvalues > | 0 = | 95 | Mardia fit | measure 1 = | 0.798 |
| Retained dime | nsions = | 3 | Mardia fit | measure 2 = | 0.993 |
| | | abs(eige | envalue) | (eigenv | alue)^2 |
| Dimension | Eigenvalue | Percent | Cumul. | Percent | Cumul |
| 1 | 964.93863 | 72.25 | 72.25 | 98.79 | 98.79 |
| 2 | 67.007783 | 5.02 | 77.27 | 0.48 | 99.2 |
| 3 | 30.236075 | 2.26 | 79.53 | 0.10 | 99.3 |
| 4 | 21.574791 | 1.62 | 81.15 | 0.05 | 99.4 |
| 5 | 15.707688 | 1.18 | 82.33 | 0.03 | 99.4 |
| 6 | 9.2522026 | 0.69 | 83.02 | 0.01 | 99.4 |
| 7 | 7.5743302 | 0.57 | 83.59 | 0.01 | 99.40 |
| 8 | 5.9310838 | 0.44 | 84.03 | 0.00 | 99.4 |
| 9 | 4.5038763 | 0.34 | 84.37 | 0.00 | 99.40 |
| 10 | 4.1936765 | 0.31 | 84.68 | soci | oloav |



Main features

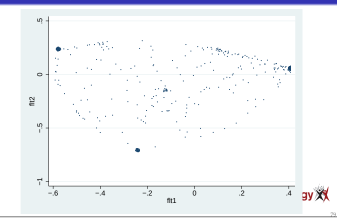
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- The homogenous clusters (1, 4, 7 and 8) are distinctly located
- Clusters with substantial amounts of more than one state are located between these "vertices" and are quite diffuse
- Some evidence of "strings" adjacent trajectories that differ slightly
- Quite clear structure, but what does it tell us about the meaningfulness of grouping? simple sequences are distinct but complex sequences are more evenly distributed

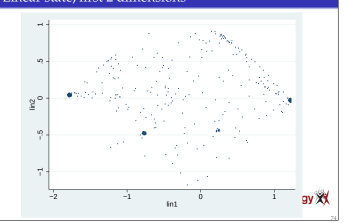
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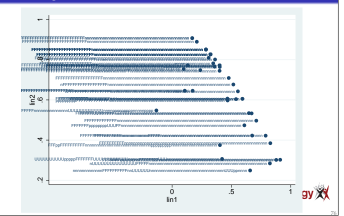






MDS of birth sequence data

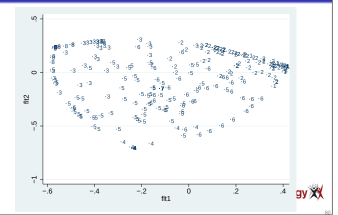
A "string" in cluster 2



Main features: dimensions

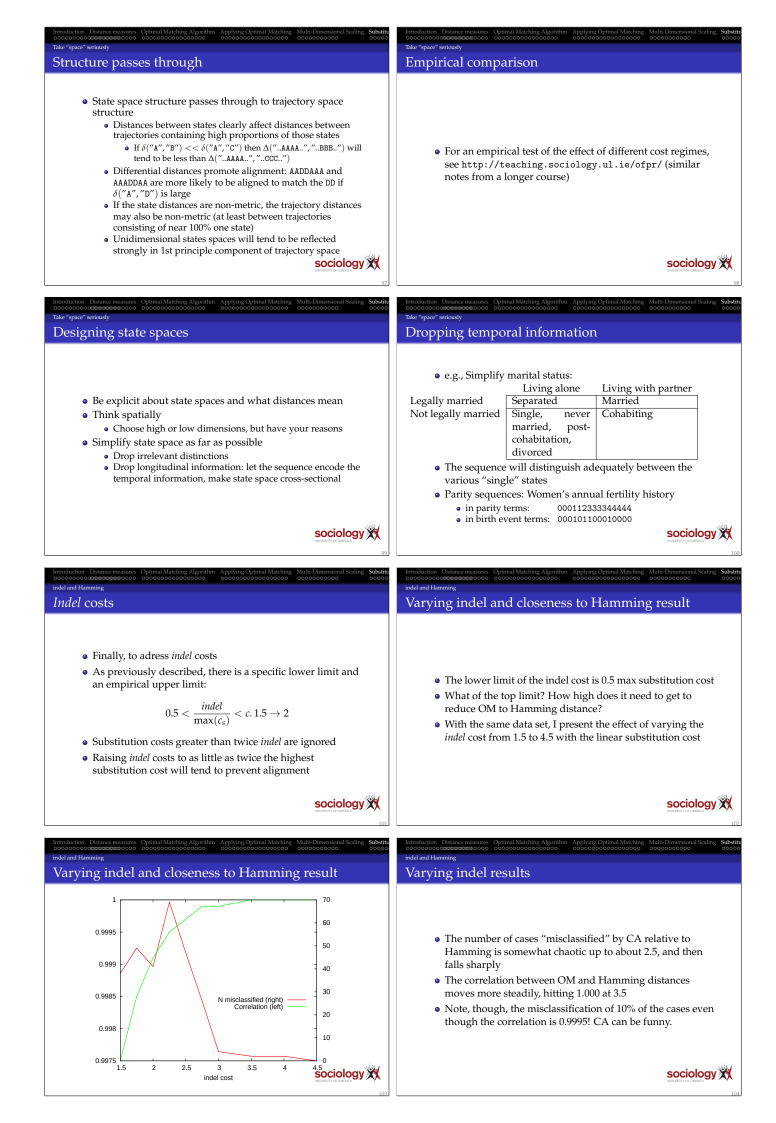
- Dimension 1 is strongly related to the dimension of the state space
 - Set FT 0, PT 1, UE 2, NonE 3
 - Sum over the time-span to give a weighted cumulative duration
 - Correlation with first dimension is 0.9999
- Dimension 2 has non-employed who become part-timers at the low end, and the transition to non-employment at the top end

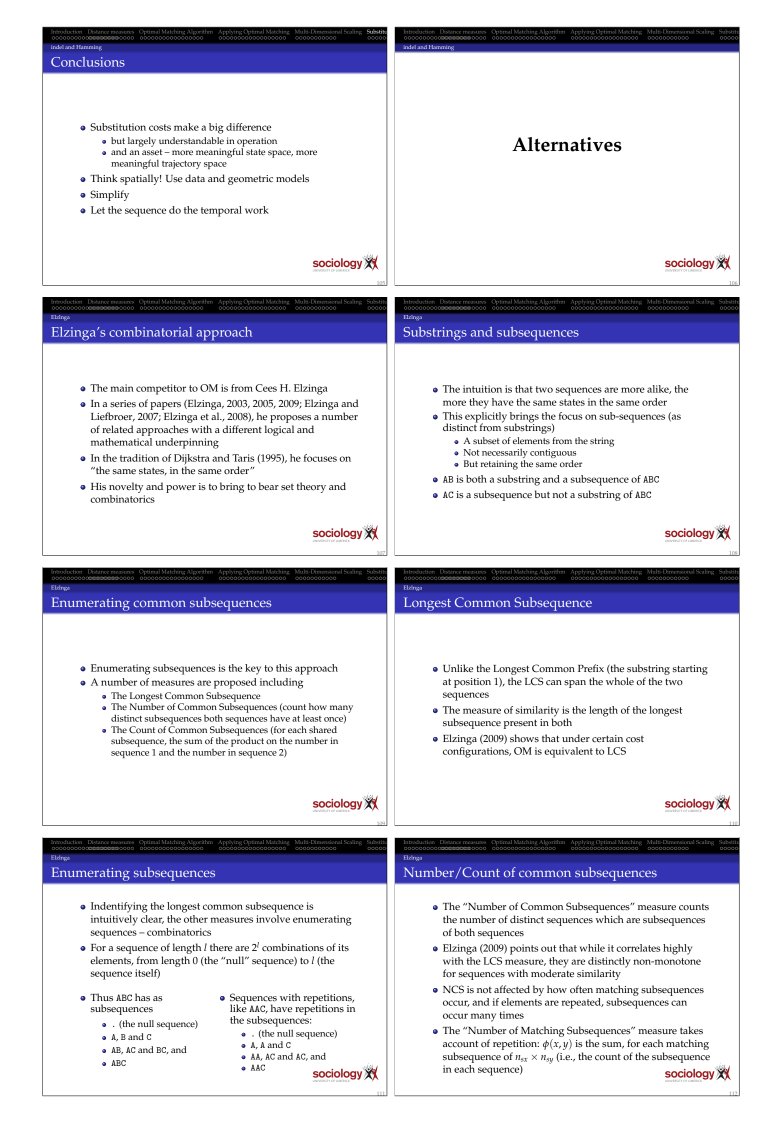




MDS useful Main features in the "flat" state space • Like before, the homogenous clusters are distinct and the As we see MDS throws light on some of the clustering others spread between them, with strings etc. processes Overall shape like a distortion of first one Distinct tight clusters of simple sequences • Dimension 1 is nearly as strongly related to the dimension • Systematically located intermediate sequences – but not in of the state space "obvious" groups Correlation 0.9947 Cluster analysis discriminates but cluster membership is • Dimension 2 runs from 100% part-time to full-timers who sensitive to small changes exit late to non-employment (100% FT also high on D2) • The main dimensions are often interpretable, and may in • Affected by the state space and the nature of the trajectories some circumstances be useful as variables in further • Trajectory state space is structured by the states, each analysis forming a pole sociology XX sociology 💥 The problem of substitution costs Does Optimal Matching make sense for sociological data? • Is the algorithm itself suitable? • How to parameterise it: substitution and indel costs Substitution and indel costs Repeated claims in the literature: that sociologists don't know how to set substitution costs, that we can't match the effectiveness of molecular biology Yes, our analytical goals are often much less well defined than those of the biologists No, substitution costs are not an intractable problem sociology 💥 sociology 💥 Criticism and puzzlement Mapping states to sequences The essence of SA is mapping a view of a state space onto a • Wu (2000) treats the choice of substitutions costs as an view of a trajectory space: $\delta(s) \rightarrow \Delta(S)$ insurmountable - while he has some misunderstandings • We start with knowledge or a view of how states relate to that made it particularly difficult for him, it is an important each other (what states are like each other, what states are stage of the OM process dissimilar) • Many other writers agonise over the problem • With a suitable algorithm we map this perspective onto • Many opt for using transition rates to get data-driven cost trajectories through the state space: what trajectories are Or set all substitution costs equal to 1 more or less similar • Neither option is really neutral - we need to understand The nature of the algorithm determines substitution cost setting better • Whether the mapping makes sense • Similarly, how to set indel costs? • Exactly how the structure of the state space affects the structure of the trajectory space sociology 💥 sociology 💥 **OMA** coherent? Thinking about state spaces and distances Costs can be thought of as distances between states • Can we expect OMA to provide a coherent $\delta(s) \rightarrow \Delta(S)$ • If state space is \mathbb{R}^n , distance is intuitive mapping? Elementary operations are intuitively appealing: If state space is categorical, how define distance? State space as efficient summary of clustered distribution in $(ABC, ADC) = f(\delta(B, D))$ (abcd, Abd) = f(indel) \mathbb{R}^n : distances are between cluster centroids State space can be mapped onto specific set of quantitative minimising concatenation of these two operations to link 3 dimensions: each state located at the vector of its mean any pair of trajectories values: Euclidean or other distances between vectors • If 3 is reasonable, 1 and 2 determine how state space affects States can be located relative to each other on theoretical trajectory space grounds sociology sociology 💥

Transitions and substitutions Informative transition rates • Transition rates frequently proposed as basis for No logical connection between substitutions and transition substitution costs rates Critics of OMA complain of substitution operations but under certain circumstances transition rates can inform implying impossible transitions (e.g., Wu, 2000) us about state distances • Even proponents of OMA are sometimes concerned about If state space is a partitioning of an unknown ℝⁿ, "impossible" transitions (e.g., Pollock, 2007), movement is random (unstructured), and the probability But substitutions are not transitions, not even a little bit! of a move is inversely related to its length, then substitutions happen across sequences, Distance between states will vary inversely with the $\Delta(ABC, ADC) = f(\delta(B, D))$ (similarity of states) transition rates • transitions happen within sequences (movement between However, these conditions are often not met states) sociology 💥 sociology 💥 Confusing state and trajectory information Deceptive transiton rates This procedure confuses party state space and voter • Example: using voting intentions as a way of defining characteristics inter party distances Voter polarisation/loyalty is trajectory information, not • UK: relatively high Con-LibDem two-way flows; ditto state information Lab-LibDem • There is a strong analytical argument for trying to keep the • But Con-Lab transitions much lower: implies a potentially two concepts as separate as possible incoherent space (non-metric, more below) • $\delta(\text{Con, Lab}) > \delta(\text{Con, LibDem}) + \delta(\text{LibDem, Lab})$ Another type of problem: irrelevant distinctions can cause similar states to have low transition rates sociology 💥 sociology 💥 Take "space" seriously Looking at state spaces • Two very simple state spaces: • Single dimension, equally spaced: • Very useful to think in spatial terms State space as efficient summary of clustered distribution in 1 2 \mathbb{R}^{n} 2 1 State space mapped onto specific set of quantitative 1 0 1 dimensions 2 1 0 State space defined on theoretical grounds • For 1 and 2, explicitly multidimensional, in case 2 All states equidistant − n − 1 dimensions dimensions are explicit • For 1 and 3, we can attempt to recover the implicit 1 0 1 dimensions 1 1 0 1 1 1 0 sociology 💥 sociology 💥 More dimensions 2-D example D • E.g., 2D picture of inter-party distances: location on left-right scale, plus on pro-/anti-EU scale • Distances are Euclidean or other metric (e.g., L1) • Euclidean: $\sqrt{\sum_i (r_i - s_i)^2}$ sqrt(5) -• L1 (city block): $\sum_i |r_i - s_i|$ · Generalises easily to many dimensions • Problem: how to weight different dimensions? • Scale by standard deviation? Substantive importance? А в С sociology 💥 sociology 💥





A distance from similarity Algorithms • As mentioned before, Elzinga (2009) asserts that where • The problem with subsequences is that there are 2^l of them – enumerating them is $O(2^N)$ $\phi(x, y)$ measures the amount of a characteristic shared by x and *y*, and the following holds: • Elzinga 2005 outlines an algorithm to enumerate common subsequences in a pair of sequences that is $O(l_1 \times l_2)$ $\phi(x,y) = \phi(y,x)$ • This is implemented in CHESA, downloadable from http://home.fsw.vu.nl/ch.elzinga/ $0 \le \phi(x, y) \le \min \left\{ \phi(x, x), \phi(y, y) \right\}$ • I have implemented a "brute-force" algorithm for Stata, then which enumerates subsequences in a first pass, and then $d(x,y) = \phi(x,x) + \phi(y,y) - 2\phi(x,y)$ compares them in a rapid second pass - good for up to is metric about 20 tokens in its current version sociology 💥 sociology 💥 Example with birth/labour market sequences Comparison with OM First problem: 73 periods is far too many (for my • The algorithm is the main difference from OM, but algorithm at least) • The absence of substitution costs is another important • Solution: sample every 4th month to yield 19 tokens per 5 difference - states are either the same (1) or different (0) year sequence • Elzinga (personal communication) has outlined a measure Number of matching sequence measure calculated and which takes account of partial as well as complete distance as similarity • But what does the NMS measure look like in practice? $d(x,y) = \phi(x,x) + \phi(y,y) - 2\phi(x,y)$ sociology 💥 sociology 💥 Cluster analysis Eight-cluster solution Full time employed • Cluster analysis on the pairwise distances yields the Part time employed following comparison with OM on the same sequences 100 Unemployed 200 (8-cluster solutions) Non-employed 300 OM NMS 1 4 5 8 100 1 | 252 0 0 0 0 9 0 0 200 2 0 27 0 0 0 51 0 0 300 3 0 0 19 0 0 23 0 0 0 5 10 15 20 4 | 0 0 0 50 0 19 0 0 10 0 38 27 5 6 0 0 3 0 0 0 0 0 0 0 0 0 0 100 7 2 0 0 0 0 0 15 0 8 0 0 0 0 2 127 200 1 0 300 0 5 10 15 20 0 5 10 15 20 sociology 💥 gy 💥 Graphs by ne8 The "interesting" clusters MDS solution: first two dimensions ڡ Full time employed 0 Part time employed Unemployed 50 Non-employed 100 150 ·6.1.66 50 2 100

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Graphs by ne8

10 15 20 0 5 10 15 20

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| Is OM optimal for life course data? | Spells as sequences of tokens |
| Lifecourse data is usually spell structured – a sequence of periods in a single state, with a given duration How to deal with in OM, which works with sequences of tokens? Treat spells as tokens, ignore duration? Represent time by multiplying tokens by spell duration? | The latter approach is usual, but it this sociologically optimal? For instance, OM says AAAB is as distant from AACB as from AABB (given δ(A, B) = δ(A, C)) Substantively, the first and third are very similar while the second introduces a completely new spell Do we need an algorithm that is aware of spells? |
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| A simple but extreme strategy is to ignore duration Makes sequences of spells, ignoring length It works - the sequences are still sequences, but now are unequal in length But duration is important: FFnnnnnn and FFFnnnnnn are substantively closer to each other than to FFFFFFFnn but all reduce identically to Fn as spell sequences | Another approach is to represent spells non-linearly Represent spell duration as e.g., √l Thus the "cost" of deleting a unit is bigger in a short spell (the fixed cost of deleting a unit represents more time in a long spell) One way of implementing this is OMv, a relatively simple adaptation of the OM algorithm (Halpin, 2010) |
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| This approach means that the cost of edits to the token strings are sensitive to the length of the spell the token is in It produces distances not very different from OM unless there is a very high level of variation in spell length However, as currently implemented it is not a stable solution Sequences composed of few, long spells are judged closer to all other sequences Making for non-metric distances Potential solutions in scaling distances according to the most-dissimilar possible sequence | Matissa Hollister proposes a similar approach (Hollister, 2009) She costs <i>indels</i> differentially according to the element's neighbours To insert z between i and j: C_{izj} = x ⋅ w_{max} + y ⋅ w_{zi} + w_{zj} where w is substitution cost This likely has similar consequences to OMv |
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| (Lesnard, 2010) has proposed a "dynamic Hamming" distance, suitable for where time has a clear scale (daily, weekly, annual) Distances are inversely related to the moment-by-moment transition rate Thus states are closer at moments when many changes happen Further apart in times of low change Successfully used with daily time-use data | OMv "warps time" by weighting it differently in different spells Harks back to Abbott's use of the term to suggest non-linear time scales (Abbott and Hrycak, 1990) In turn informed by Sankoff and Kruskal (1983), <i>Time Warps, String Edits and Macromolecules</i> |
| | |

| Time "warping" | 00000000000000000000000000000000000000 |
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| Time warping algorithms | Multiple domains |
| Formally, time warping is a family of algorithms that do "continuous time-series to time-series correction" while OM <i>et al</i> do "string to string correction" (Marteau, 2007) Focus on comparing pairs of continuous-time high-dimensional time-series in ℝⁿ Operates by locally compressing or expanding the time scale of one trajectory to minimise the distance to the other Distance is usually Euclidean in ℝⁿ or other simple distance | Handling trajectories through multiple domains simultaneously is very attractive Quite a few examples in the literature, often combining Labour market Housing Partnership and family formation Dijkstra and Taris (1995) use as an example residential, educational and job status Pollock (2007) uses a similar trio |
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| Multiple domains | Multiple domains |
| Multiple state spaces | Substitution costs in multiple spaces |
| The immediate difficulty is how to deal with multiple state spaces One solution is to create a combined space, crosstabulating the others, e.g.,: employed-single employed-partnered not employed-partnered not employed-partnered However, in practice this usually generates a high number of cells Practical problem of determining substitutions costs | It may be possible to set the costs on the cross-tabulated spaces <i>a priori</i>, using intuition or theory Sometimes it may also be possible to simplify the structure of this space: e.g., collapse certain regions into a single category If clear state-space structures exist for the sub-spaces it may be possible to combine them systematically: Euclidean: √∑_i(x_{ij} - x_{ik})² Sum the different distances: ∑_i x_{ij} - x_{ik} Weighting subdomains differentially is also possible |
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| COCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCO | Multiple domains Multi-channel SA |
| | |
| Parallel analyses An other option is to conduct parallel analyses in each domain This yields multiple distance measures Allows analysis of how the different domains cluster independently Perhaps less sensitive to coordination issues within lifecourses but should still be interesting | Multi-channel SA A team in Switzerland are drawing on newer bioformatics technology: Multi-channel SA Explicitly deals with multiple parallel trajectories Gauthier et al. (2008) describe their "MCSA" method and claim it is superior to parallel OM analyses, and simpler than handling multiple-state distances Software available: http://www.tcoffee.org/saltt Not entirely clear from the description how the method works Bühlmann (2008) uses their methodology to examine careers of Swiss economists and engineers, using a number of categorical measures of their status |
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Stata and OM

• Kohler et al provide the SQ add-on for Stata:

. ssc install sq

TraMineR

- TraMiner (http://mephisto.unige.ch/traminer) is an R package for sequence analysis
- R is a free/open-source implementation of the S-Plus ge(http://www.r-project.org)
- ble for most platforms (including Windows and
- owerful, good graphics, but R is not ner-friendly

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