# Accelerating Biomolecular Nuclear Magnetic Resonance Assignment with A\*

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## Overview

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Motivation

Introduction

#### Motivation

- Nuclear Magnetic Resonance Spectroscopy
  - Gain knowledge about protein structure
  - Study how mutations lead to diseases
- Problems
  - Generates large amounts of data
  - Data analysis is slow and error prone
- Goal
  - Automate the assignment process
  - Decrease human error
  - Increase productivity

Nuclear Magnetic Resonance Spectroscopy

Introduction •0

# Nuclear Magnetic Resonance (NMR)

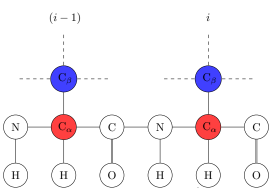
- Used to obtain structural information
  - Chemical shift values
- HNCACB experiment
  - Generates  $C_{\alpha}$  and  $C_{\beta}$  residue i and i-1
- CBCA(CO) NH experiment
  - Generates  $C_{\alpha}$  and  $C_{\beta}$  for residue i
  - Confirms residue data

Nuclear Magnetic Resonance Spectroscopy

Introduction 0

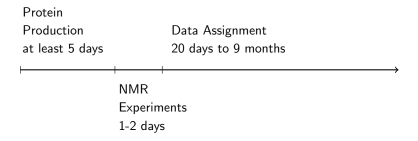
## Chemical Shift Values

## HNCACB



Data Collection and Manual Assignment

#### Timeline



Data Collection and Manual Assignment

#### Manual Methods

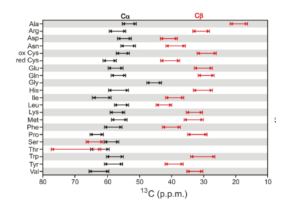
- Most time consuming part
- Prone to human error
- Missing and ambiguous data forces chunks to be skipped

## Initialization

- Input
  - Expected amino acid sequence
    - Converted to expected chemical shift values
    - Stored as the reference protein chain
  - NMR chemical shift data
    - $C_{\alpha}$  and  $C_{\beta}$  for residue i and i-1
    - Stored in a tile
- Missing data
  - Place holder tile generation
- Grouping

Preprocessing

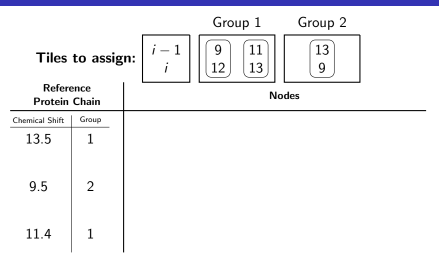
# Grouping



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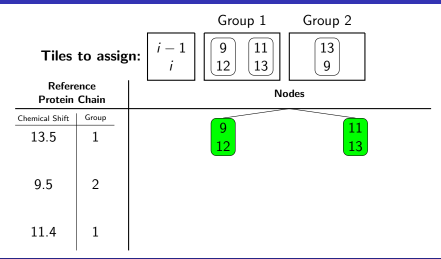
Assignment

# Starting the assignment



Assignment

# Starting the assignment



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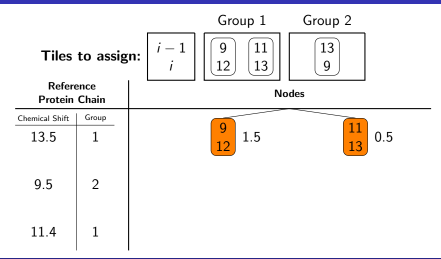
Assignment

#### Cost Calculation

- Accuracy matching the protein chain residue
- Accuracy matching the tile above current tile
- Cost of all tiles place before current tile

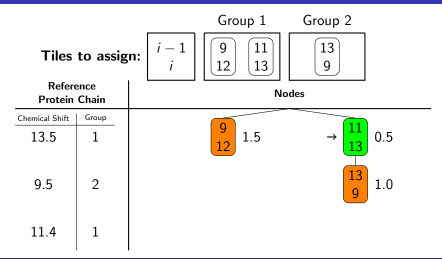
Assignment

# Generating child nodes



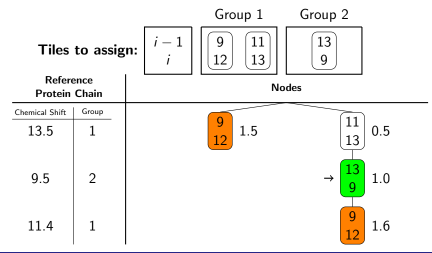
Assignment

## Generating child nodes



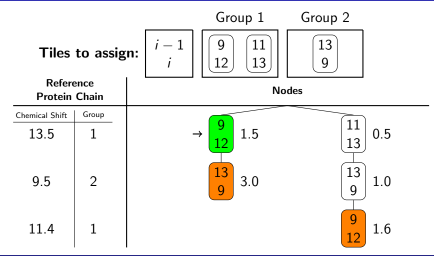
Goal State

## Goal State



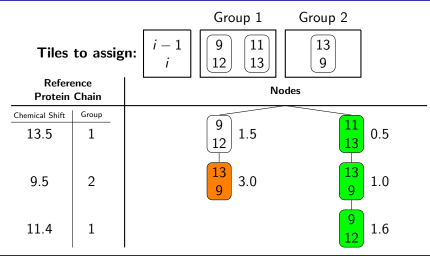
Goal State

## Goal State



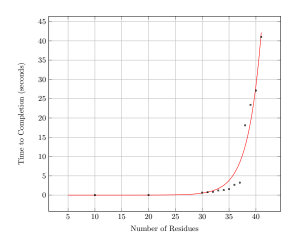
Goal State

## Solution State



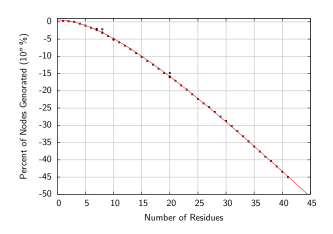
Results

# Time of Assignment



Results

#### Child Nodes Genorated



Outlook

#### Future Goals

- Parallelization
  - Decrease assignment time
  - Allow for larger data sets
- Machine learning
  - Increase accuracy of assignment
  - Optimize cost calculation

Outlook

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- Leah Robison (research colleague)

Bibliography

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## Thank You

