# Bayesian Analysis of Esports Teams' Perfomances Alto University

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## 1 Introduction

In this report, we present a Bayesian Analysis approach for evaluating team performances for the esport *Dota* 2. This topic was chosen due to the personal interest of our team members in esports. We formed the group based on this and all of us were familiar with either Dota 2 or League of Legends.

Before describing the problem more thouroughly, we would like to provide a brief insight into what we are trying to achieve. Dota 2 is a MOBA game and an esports discipline. Esports have a similar structure as traditional sports, where there are professional seasons which roughly equal to a year, and these are finished with a main tournament (in our case: *The International (TI)*).

The tournament itself consists of two stages: qualifications and main event. The latter is also split into two parts: group stage, and play-offs. During the group stage, teams compete in order to have a better starting position in the play-off stage. For our analysis, we used the data from the last TI where the best teams in the world competed.

The professional Dota 2 scene is divided into different regions (e.g. Europe, CIS, China, etc). Teams from these regions participate in qualifications separately, and the very first goal of the team is to become the best within their region. As a result, teams from one region play with each other much more often, and it can influence the way they perform. Hence, we also used information about the region of the teams while designing and implementing our models.

Our aim is to find a way to predict the match result based on the performances of the competing teams (the way performances are computed is discussed later). In other words, we propose a way to make the analysis of the previous matches in which teams participated, and take those results to predict how they would perform in the future.

# 2 Data and Preprocessing

In our analysis, we decided to use matches from **The International 2019** Tournament, the results of which are available online through some APIs. The data was gathered with the Python programming language in two steps. At first, match IDs were collected from DotaBuff. Afterwards, we used the obtained IDs in order to acquire detailed information about each match. For the latter we used the OpenDota API.

For our analysis, we used the data from the group stage of the tournament as each team played the same amount of matches. On the other hand, for the posterior predictive checks (which are further discussed in the Posterior Predictive Checks section), we made predictions for the play-off stage of TI9.

The next step was to define a new metric in order to quantify the teams' performances. To do this we used a mixture of different measures/scores which we then normalized and used its sum. The initial features can be seen below:

Variable Names	Description
radiant_score	Final score for the Radiant team(number of kills on Radiant)
dire_score	Final score for the Dire team (number of kills on Dire)
radiant_xp_adv	Array of the Radiant experience advantage at each
	minute in the game. A negative number means that
	Radiant is behind, and thus it is their experience disadvantage.
radiant_gold_adv	Array of the Radiant gold advantage at each minute in the game.
-	A negative number means that Radiant is behind, and thus it is
	their gold disadvantage.
hero_damage	Hero Damage Dealt (user specific)
hero_healing	Hero Healing Done (user specific)
obs_placed	Total number of observer wards placed (user specific)
kda	kda (ratio of kills/deaths/assists) (user specific)

These features were extracted for each match and contain information about both of the teams. Since each *Dota2* game consists of 5 players, this means that we end up with 10 data points for each match.

As you can see from the table above, the first 4 features have to do with team-wide statistics while the rest 4 have to do with user-specific statistics since they evaluate the performance of each player separately. Our goal is to find a way to combine these features so that the result will be a performance score for each team, for each match.

Our approach to this can be described by the following equation which assigns a score to each team for each match:

$$R_{score} = score\_gap + exp\_advantage + gold\_advantage + \\ \sum_{\forall p \in P_R} hero\_damage(p) + healing\_done(p) + kda(p) + wards\_placed(p)$$

where  $R_{score}$  denotes the radiant team score, and  $P_R$  is the set of all radiant players on the specific match. The same was also applied for the dire team.

NOTE: This equation is over-simplified since it does not take normalization into account (having all values in the same range).

Each of these functions is described below:

Variable Names	Description
score_gap	Radiant score minus Dire score (so it can also be negative).
exp_advantage	A weighted sum of the experience advantage that Radiant had during
	the game (can also be negative). We are giving bigger weights
	to the scores at the end of the game since they are more
	informative about the final result.
gold_advantage	A weighted sum of the gold advantage that Radiant had during
	the game (can also be negative). We are giving bigger weights
	to the scores at the end of the game since they are more
	informative about the final result.
hero_damage	Simplistic Approach: The difference of the sum of the damage dealt by
_ 0	all Radiant players when compared to Dire players.
healing_done	Simplistic Approach: The difference of the sum of the healing that
	was done by all Radiant players when compared to Dire players.
kda	Simplistic Approach: The difference of the sum of kdas of
	all Radiant players when compared to Dire players.
$wards\_placed$	Simplistic Approach: The difference of the sum of the number of
	observation wards that were placed by all Radiant players when
	compared to Dire players.

As you can see, a rather simplistic approach was used to combine the data. As a normalization technique we are currently using MinMax scaling which does not completely distort our values and so it will keep the following notion:

- Large performance scores should probably result in a Radiant win (not always).
- Smaller scores (negative ones) should point us to the direction that Dire will most probably win.

For more information about the implementation you may check the data transformation appendix. An important note here is that, for each match, the  $score\_gap$ ,  $exp\_gap$  and  $gold\_advantage$  columns for the Dire team are multiplied by -1 so that they will be positive in case the Dire has positive statistics (and negative otherwise).

# 3 Problem Solving

By preprocessing our data we managed to get a single scoring measurement for each team. Our next step is to build a model on top of these transformed data and try to predict/sample future scores (future performances). Our approach involved two models which are going to be described below. For the implementation we used the RStan (Stan Development Team 2020) package of R.

At first, we are going to represent our data as a 2-dimensional array which will contain information about each team. The rows will denote the matches and the columns will denote the teams, while the content of each array cell will be the above computed performance scores.

Below, we are going to describe two model-architecture approaches, hierarchical and separate models. In the first case, under the hierarchical normal model, we assume that our data (Dota 2 matches per team for each region) are independently normally distributed within a certain number of groups (which is equal to the number of teams that each region has). Here, all groups share a common variance  $\sigma^2$  while they have different (team-specific) means  $\theta_j$  (Gelman et al. 2013). We also assume that the means follow a certain distribution (disccussed in the next section) with unknown mean  $\theta_{\mu}$  and standard deviation  $\theta_{\sigma}$ .

In the case of separate models, we assume that each team has its own model and so we will build separate models which will contain no other information other than the performances of the teams on their matches. To implement that, we use separate priors  $\theta_i$  and  $\sigma_i$  for each of the teams.

#### 3.1 Choice of Priors

Our first trials consisted of trivial priors which contain no problem-specific information, meaning that we did not use any current knowledge while picking them. The reason we chose these weakly informative priors is because we did not want to use current knowledge since it would skew our results towards a certain assumption (e.g. that the teams of region X perform better) which would not hold in the general case.

While tuning these priors we noticed that the results and the output diagnostics were mostly the same. This made us stay with the choices above since these weakly informative priors showed some stability.

In our problem we represent our observed performances by  $y_{ij}$ , where i = 1..N denotes the current match and j corresponds to the team of a certain region. So  $y_{ij}$  is the performance score of team j in game i. We can now mathematically express the separate model as:

$$y_{ij} \sim N(\mu_j, \sigma_j)$$
  
 $\mu_j \sim N(0, 100)$   
 $\sigma_j \sim Inv - \chi^2(0.1)$ 

For the hierarchical model these weakly informative priors can be mathematically expressed as follows:

$$\theta_{\mu} \sim N(0, 100)$$

$$\theta_{\sigma} \sim inv - \chi^{2}(0.1)$$

$$\theta_{j} \sim N(\theta_{\mu}, \theta_{\sigma})$$

$$\sigma \sim inv - \chi^{2}(0.1)$$

 $y_{ij} \sim N(\theta_i, \sigma)$ 

In the representations  $\theta_{\mu}$  and  $\theta_{\sigma}$  are the hyper-priors of the means (and the means are denoted as  $\theta_{j}$  for each team).

## 3.2 Separate Model

Firstly, we built a separate model which could reflect each team performance separately. The idea behind the separate model is that all teams are unique and there is nothing common with them. This contradicts with our hypothesis that teams from different regions tend to have different gameplays and strategies (e.g. teams from region X might follow a more offensive gameplay).

In the separate model, each team is going to have their own mean and variance. This means that, when we later reach the prediction stage, we cannot make any predictions about unseen teams (this is not a problem in our case) and we can also not extract any generic information about the gameplay and the strategies.

#### 3.2.1 Using the model

You may find the stan implementation of the separate model in the appendix (Stan Code). Now in order to run the separate model on our dataset we are using the following R code:

From the above code you can see that we used the default configurations for compiling stan models with the exception of the chains parameter which is set to 5 instead of the default 4. This means that we are running 5 chains and for each chain we do 2000 iterations and we prune the first 1000 (warm-up iterations) since these are the default values.

#### 3.3 Hierarchical Model

The goal here is to create a hierarchical model for the teams of each region. As we have already mentioned above, we can divide the 18 teams to six regions (Europe, China, CIS, Southeast Asia, North America, South America). By doing that we get different posterior samples for the teams of each region and this can later help us define how different is the gameplay and the strategies used in each region.

Now, in order to build the models, we are going to use the same hierarchical structure for all different regions. This way we are sure that there is no external bias.

#### 3.3.1 Using the Model

You may find the stan implementation of the hierarchical model in the appendix (Stan Code). Here, for each region, we run the hierarchical model on its data set points (the matches of the teams of that region). For example, in the Europe region, we have the team Alliance, CHAOS, OG, NiP, Secret, Liquid to form the data set.

```
chains=5, seed=42)
draws_Europe_hm <- as.data.frame(model_Europe_hm)</pre>
```

Again, you can see that we used the default parameters of stan and that is true for every hierarchical model of every region.

# 4 Convergence Diagnostics

Our basic method of checking for convergence was using the  $\hat{R}$  diagnostic. For the implementation we used rstan's (Stan Development Team 2020) Rhat function which compares the between and the withing-chain estimates of the model parameters and other univariate quantities of interest.

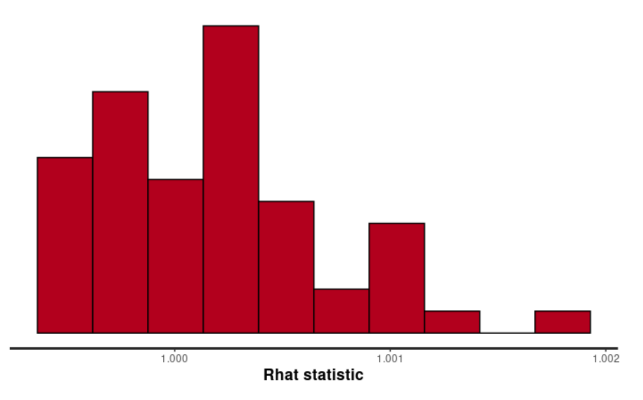
An  $\hat{R}$  value larger than 1 means that the chains have not mixed well and that they don't converge to a common distribution (so the between and within-chain estimates don't agree). On the other hand, values really close to 1 are good indicators of convergence of the chains.

## 4.1 $\hat{R}$ Results

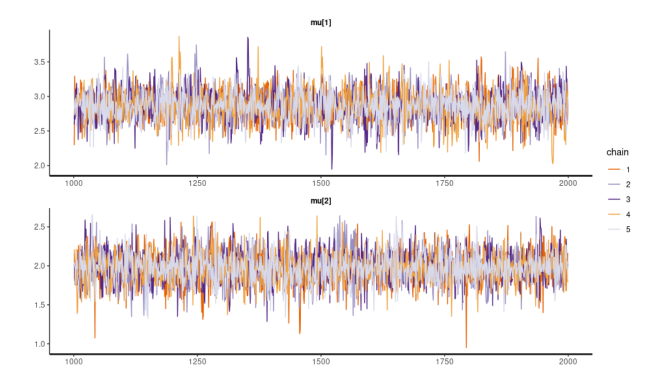
#### 4.1.1 Separate Model

For the separate model we ended up with and  $\hat{R}$  value of 1 for all of the parameters (these include 18  $\mu$  parameters and 18  $\sigma$  parameters, one for each team). This implies that our chains converge to a common distribution which is what we aimed for.

The  $\hat{R}$  plot below shows the values that  $\hat{R}$  has taken for the separate model:



As you can see, the largest Rhat value is still smaller than 1.002. In addition, in the figure below you can also see how the values of the posterior samples for  $\theta_1$  and  $\theta_2$  are evolving (after removing the warm-up iterations).



#### 4.1.2 Hierarchical Model

With the above weakly informative priors we managed to reach converge after compiling 5 stan models (one for each region). The  $\hat{R}$  values have a similar distribution as above. In particular, all of the  $\hat{R}$  values (rounded to 3 decimals) had a value of 1 which means that we reached convergence again.

Here, it is worth mentioning that the North American region has only one team participating the TI which means that only one team defines a whole region. Ideally we would need more information about the region before performing the modeling.

## 4.2 Effective Sample Size

Since the number of posterior parameters is too large, we cannot include the effective sample size for each one of them. In general, an effective sample size (ESS) greater than 100 (per chain) is considered good. In our case, we used the Bulk and Tail ESSs and we got values close to 5000 for each parameters for both the separate model and the hierarchical ones. This is a good indicator that our posterior estimates are reliable.

#### 5 Posterior Predictive Checks

We compare the MCMC results with the performance data which we used.

# 6 Model Comparison

## 7 Predictive Performance Assessment

#### 7.1 Predictive Performance

The prediction workflow for a series match between team X and Y is the following:

- 1. A normal series is a two out of three game series. But here we only compare each team's expected performance scores to predict who is more likely to win in the series.
- 2. The *ypred* in the stan code represents the predictive performance score of each team. So, by comparing the *ypred* of team X and Y, we could get the win possibility. The function predict\_result\_sm below will do this:

```
predict_result_sm <- function(team_num_1, team_num_2) {
    ypred_1 <- draws_sm[, 36 + team_num_1]
    ypred_2 <- draws_sm[, 36 + team_num_2]
    win_rate_1 = {}
    for (i in 1:length(ypred_1)) {
        win_rate_1[i] = ypred_1[i] / (ypred_1[i] + ypred_2[i])
    }
    mean_rate_1 <- (mean(win_rate_1))
    return(mean_rate_1)
}</pre>
```

3. If the win possibility of team X is above 0.5, we will say that the team X is more likely to win in this game series.

Here is an example to predict which team will win between PSG.LGD and VP.

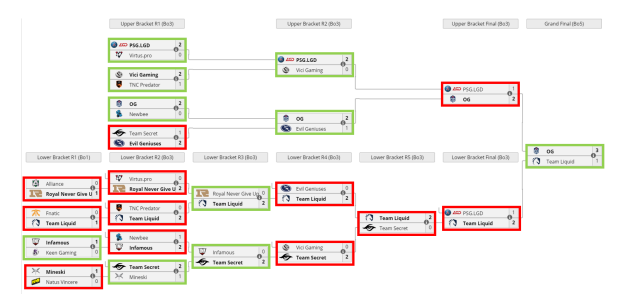
```
#return the win team
compare <- function(team_name_1,team_name_2){
  team_num_1 <- which(colnames(data) == team_name_1)
  team_num_2 <- which(colnames(data) == team_name_2)
  predict <- predict_result_sm(team_num_1,team_num_2)
  if(predict>=0.5)
    return(team_name_1)
  else
    return(team_name_2)
}

#Calculate the result of main event stage
compare("PSG.LGD","VP")
```

#### ## [1] "PSG.LGD"

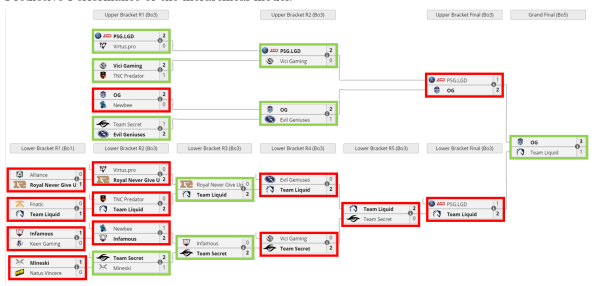
We actually apply this method to all of the series in the main event. Here are the prediction result compared to the real ones. The green boxes indicate that the predictions correspond to the actual results, while red boxes indicate that the predictions do not correspond to the actual results (missclassification).

Predictive Performance of the separate model:



From the above image we can see that we reached an accuracy of 45.5% with the separate model.

Predictive Performance of the hierarchical model:



Here we can see that the accuracy is almost 41% which is lower than the separate model.

## 7.2 Analysis

As you can see, there is no significant difference in the results between the separate model and the hierarchical model. This proves that regional factors do not play a significant role. In other words, there are no significant differences between the teams in the different regions.

It is also worth mentioning that the prediction accuracy is not very high. Possible reasons are the following:

1. We just predict the main event result based on group stage performance. It often happens that a team is in poor form in the group stage, but gets better and better in the knockout stages (just like in normal sports). With this type of teams, our predictions are difficult to be accurate. For example, Team Liquid was in the 14th spot out of all teams in the group stage, but in the end it finished at second place. So we were almost entirely wrong in our predictions for this team.

2. Our performance score/metric is not ideal and does not completely reflect the power of each team. This is discussed in more detail in the improvements section.

# 8 Prior Sensitivity Analysis

As mentioned above, trying out different weakly informative priors yielded similar results and the models converged in all cases. So, in order to avoid clutter we chose not to include more results since the output figures are almost the same.

Below we are going to mention the another choice of priors which is considered more informative.

## 8.1 Separate Model

Here, we chose to use the mean performance score and its standard deviation from the gathered data as our prior for the mean and sigma of the separate model, respectively.

Mathematically this can be noted as:

$$\mu_i \sim N(\mu_0, \sigma_0)$$

where  $\mu_0$  denotes the mean performance score for the gathered data, and  $\sigma_0$  stands for standard deviation of the data.

The standard deviation for the sampling is now

$$\sigma_j \sim inv - \chi^2(\sigma_{new})$$

where  $\sigma_{new}$  is drawn from  $\mu_0 - N(\mu_0, \sigma_0)$ .

The above configuration resulted in an accuracy of 52.5% (correctly predicted 12 out of 22 matches).

#### 8.2 Hierarchical Model

For the hierarchical model, we used the following mathematical notation:

$$\theta_{\mu} \sim N(\mu_0, \sigma_0)$$

$$\theta_{\sigma} \sim inv - \chi^2(\sigma_{new})$$

where the  $\sigma_{new}$  was drawn in the same way as in the case of the separate model.

The above configuration resulted in an accuracy of 52.5% (correctly predicted 12 out of 22 matches) which is the same as with the separate model.

# 9 Discussion of Issues and Possible Improvements

From the above results, we can see that there is a large number of issues and improvements can be made in the current work.

- 1. The data from the group stage matches does not fully reflect the ability of the teams, so we should also include the previous performance of each team in the database. Getting further information about each team's performances is crucial since the whole prediction process is based on that.
  - Here, we need to be careful with what data we choose. Just like in normal sports, the team members of e-sports teams are not always the same and transfers happen from time to time.

- This means that we should also take that factor into account. A possible solution would be to completely separate the individual players' performances so that each team's performance is not tied to games on which the team is always the same, but it is tied to the games that each player of the team has played. This way each team's performance could be the sum of its players' performances irregardless of whether they play on the same team or not.
- The above is a nice way to gather more data but does not take into account the fact that teamwork is also an important part of e-sports (how players collaborate with each-other).
- 2. The method of adding up all the parameters to get a performance score is too simple. Perhaps we should find a more accurate way of representing the performance of the individual teams.
  - For example, by assigning different weights to performance indicators based on group stage ranking, so that the final score matches their actually performance.
  - Another approach would be to follow the note of the previous bullet (that teams are not always the same) and gather more data by using player-specific knowledge.

## 10 Conclusion

Above we described a way to predict future statistics about Dota 2 matches by using a very small dataset. By using Bayesian Analysis we managed to extract predictions about match outcomes and compare the gameplay of the teams of each region.

The results may not be ideal but there is room for improvement and we did not reach a dead end. In the previous paragraph we mentioned what steps could lead us to better results and how to overcome the low amount of prior information. By following these steps, we believe that we can achieve much greater accuracy and better models overall.

In addition, we saw how separate and hierarchical models are compared and discussed their strengths and weaknesses. These models helped us predict the results of the TI and the latter can also help us extract gameplay specific information. For example, by destructuring the performance scores we could see that higher performance scores mostly correspond to offensive teams and so we could use that assumption in order to predict gameplay styles for each region. We decided not to include the above since it was based on a big assumption and our data transformation could not guarantee that this holds (large performance values match to offensive teams).

In general, the above report includes a basic and straightforward approach to perform Bayesian Analysis on multidimensional data by transforming the data and using team performances as "quality" indicators. Since our prior information concluded of a very small number of matches for each team, we can say that the results are sensible while there is still room for improvement both in our data and model choices.

#### 11 Self-Reflection

#### 11.1 Rongzhi Liu

There are many things in life that cannot be accurately and simply measured by a number, such as how much he loves me. This work made me realize that it is sometimes more difficult to get the right data than it is to analyze it using Bayesian theory. In practice, there is a wide range of knowledge to be used and a variety of problems to be encountered. The practical problems often do not follow the fixed processes that we learn about on the course. For different problems, we first need to clarify the structure and requirements for solving them. If the structure is clear and precise, even a very simple model can give satisfactory results.

## 11.2 Georgios Karakasidis

We found many obstacles in our way when trying to implement the above models. At first, the data had no clear structure and so we had to find our own metric in order to quantify the performance. This was not as straightforward as one might think and we were on the verge of completely changing our project structure.

At last, we came up with the above approach which is both simplistic yet powerful enough in order to allow us to do inference.

I personally found it really interesting how a simple transformation of data can affect the results in such a way. In addition, Bayesian Analysis managed to output sensible and reasonable results even with these simplistic representations and the very low amount of data which is something very noteworthy.

# 12 Appendix

#### 12.1 Data Transformation

Below, we present the python functions that were used for data preprocessing and for computing the performance for a single team of a single match.

```
def _score_gap(match):
    # Simple score difference
    score = match[['radiant_score', 'dire_score']].values[0]
   return score[0] - score[1]
def _xp_gap(match, use_weights=True):
    # Take the sum of the xp advantages (over all minutes)
   xp_list = np.array(ast.literal_eval(pd.unique(match.radiant_xp_adv)[0]))
    # A negative sum would mean that the radiant team was on a disadvantage
    # most of the times
    if use_weights:
        # The last few minutes are more important and so we should give a
        # greater weight to these
       weights = np.linspace(0.1, 1, num=len(xp_list)) # avoid zero weights
       xp_list_weighted = weights * xp_list
        return np.sum(xp_list_weighted)
   return np.sum(xp_list)
def _gold_advantage(match, use_weights=True):
   gold_list = np.array(ast.literal_eval(pd.unique(match.radiant_gold_adv)[0]))
    # A negative sum would mean that the radiant team had less gold for
    # most of the game
    if use weights:
        weights = np.linspace(0.1, 1, num=len(gold_list))
        gold_list_weighted = weights * gold_list
       return np.sum(gold_list_weighted)
   return np.sum(xp list)
def _hero_specific_scores_v2(match): # hero damage and hero healing
    stats = {
        "damage": match['hero_damage'].values,
        "healing": match['hero_healing'].values,
        "kda": match['kda'].values,
        "wards": match['obs_placed'].values,
   } # will contain the damage and healing scores for the radiant players
    # Simply use the sum
   dmg_diff = np.sum(stats['damage'])
   heal_diff = np.sum(stats['healing'])
   kda diff = np.sum(stats['kda'])
   wards_diff = np.sum(stats['wards'])
   return dmg_diff, heal_diff, kda_diff, wards_diff
```

```
def performance(match):
    # match should be a dataframe that contains 5 rows
    # (a match entry for a certain team)
    score_gap = _score_gap(match)
    xp_gap = _xp_gap(match, use_weights=True)
    gold_adv = _gold_advantage(match, use_weights=True)
    # case 3 (simplistic approach)
    total_dmg, total_heal, total_kda, total_wards = _hero_specific_scores_v2(match)
    return score_gap, xp_gap, gold_adv, total_dmg, total_heal, total_kda, total_wards
```

By using the performance function above for both teams of every match we can get a new dataset which will contain all of the 7 characteristics returned by *performance*. We can then normalize these values with the method mentioned in the Data and Preprocessing section and then simply sum the values of each row in order to get the final performance score.

#### 12.2 Stan code

#### 12.2.1 Separate Model Implementation

```
data {
  int < lower =0> N;//the number of matches
  int < lower =0> J;//the number of teams
  vector [J] y[N];//the performance score
parameters {
 vector [J] mu;
  vector < lower =0 >[J] sigma ;
model {
  for (j in 1:J){
   mu[j] ~ normal (0, 10);
    sigma [j] ~ inv_chi_square (0.1);
  }
  // likelihood
  for (j in 1:J)
   y[,j] ~ normal (mu[j], sigma [j]);
generated quantities {
  vector [J] ypred;
  // Compute predictive performance for each team
  for (j in 1:J)
   ypred[j] = normal_rng (mu [j] , sigma [j]);
}
12.2.2 Hierarchical Model Implementation
data {
  int < lower =0> N;//the number of matches
  int < lower =0> J;//the number of teams in this region
  vector [J] y[N];//the performance score
```

```
}
parameters {
  real mu_p;
  real < lower =0 > sigma_p;
 vector [J] theta;
 real < lower =0 > sigma ;
}
model {
  // priors
  mu_p ~ normal (0, 10);
  sigma_p ~ inv_chi_square (0.1);
  for (j in 1:J){
    theta[j] ~ normal (mu_p, sigma_p);
  sigma ~ inv_chi_square (0.1);
  // likelihood
  for (j in 1:J)
    y[,j] ~ normal (theta[j], sigma);
}
generated quantities {
  vector [J] ypred;
  // Compute predictive performance for each team in this region
 for (j in 1:J)
    ypred[j] = normal_rng (theta[j] , sigma);
}
```

## References

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Stan Development Team. 2020. "RStan: The R Interface to Stan." http://mc-stan.org/.