INFORMATION GAIN OF STRUCTURED MEDICAL DIAGNOSTIC TESTS
Integration of Bayesian Networks and Ontologies

Marin Prcela, Dragan Gamberger, Tomislav Šmuc
Rudjer Boskovic Institute, Bijenicka 54, Croatia

Nikola Bogunović
Faculty of Electrical Engineering and Computing, University of Zagreb, Unska 3, Zagreb, Croatia

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Abstract: Usage of Bayesian networks in medical decision support system is in general case twofold: (1) for obtaining probabilities of occurrence of medical events (i.e. possible diagnosis) and (2) for obtaining information gain of actions that can be taken (i.e. diagnostic tests). On the other hand, typical role of ontology is to provide a framework for definition of medical concepts, their structure and relations among them. In medical practice diagnostic tests are commonly comprised of number of measurements or sub-tests – a structure which is straightforwardly described by ontological language. In this paper we are analyzing the information gain of such structured medical diagnostic tests. The purpose of this analysis is to allow finding (1) which structured medical diagnostic test is at the given point the most informative one and (2) which elementary measurements within a given diagnostic test are the most informative ones. Furthermore, we are analyzing some computational issues which arise in the reasoning process.

1 INTRODUCTION

Bayesian networks (BN) have already demonstrated their practical value in medical decision support systems. The most exploited features of such system are (1) finding probabilities of possible events in the system (usually probabilities of diagnosis) and (2) finding information gain (IG) of possible actions that can be taken (usually medical diagnostic tests). On the other hand, ontologies have become de facto standard in medical decision support systems for formalization of descriptive medical knowledge: defining domain concepts and relations among them.

Medical domain is particularly suitable for usage of IG as a decision support parameter. For example, (Jagt 2002) uses BN to describe probabilistic relations among medical concepts and uses IG to find the most informative medical measurement (test) considering possible final diagnoses (diseases). In practice, medical diagnostic tests are commonly comprised of more than one diagnostic parameters, e.g. laboratory analysis of blood sample measures levels of glucose, creatinine, cholesterol, urea, etc. It would be useful to allow the decision support system to perceive such structured medical test as a conceptual unit. Still, one should be aware that it is not necessary to measure all existing parameters within test; one can choose which parameters are currently interesting and disregard the others and thus presumably cut the expenses of medical test and save some time.

In this paper we are proposing approach which uses BN for description of probabilistic relations among medical concepts and measures IG of composite medical diagnostic tests defined within ontology. As we will demonstrate, the integration of those two knowledge formalisms brings in some additional features for the decision making but also rises some computational issues in the reasoning process.

It should also be noted that in medical practice very often some other (non-medical) factors must be taken into account: e.g. price of the test, availability...
of medical instruments, urgency, etc. Although measure of IG does not take into account these factors, it is possible to derive a weighted combination (or some other type) of those factors to form a comprehensive scale of medical test utility. However this paper analyzes solely IG of actions in pure medical sense.

The organization of paper is as follows. Chapter 2 previews the existing approaches for integration of ontologies and BN. Chapter 3 demonstrates the usage of BN in a medical decision support system. Chapter 4 upgrades the described decision support services with ontological knowledge and analyzes means for performing reasoning tasks. Chapter 5 gives example of practical usage of such integrated knowledge base in a single decision support system. Chapter 6 discusses some performance issues of the described approach related to reasoning phase.

2 RELATED WORK

Integration of ontologies with BN is not a new paradigm. In (Pan 2005) BN is used to recognize semantic relations between concepts in two different ontologies, which enables automatic generation of mappings between ontology concepts. Application of this approach is described in the domain of Semantic Web where problem of semantic relation between ontologies is emphasized.

In (Devitt 2006) knowledge stored in the ontology is used for generating possible structures of BN. Since ontologies thoroughly define domain concepts and existing relations among them there is a possibility to use such knowledge to generate the structure of BN. In (Town 2004) ontology is used both to learn BN structure and in the process of network training, i.e. learning probability tables of network nodes. The usual process of BN training (using existing data set) is augmented by scoring scheme which is based on the ontological knowledge.

In (McGarry 2007) high level knowledge obtained from ontologies is integrated with newly discovered knowledge extracted from BN which was trained on existing data set. In (Huhns 2007) ontology is used for management of evidence in the BN. In (Wang 2008) ontology is used for integration of heterogeneous data sources and BN is used for making probabilistic suggestions.

In medical domain, (Jeon 2007) uses ontology for semi-automatic construction of BN for diagnosing diseases. In (Zheng 2005) guideline modelling tool that uses ontological workflow management (GLIF) is integrated with probabilities obtained by BN.

As we have demonstrated in this brief overview, previous attempts of integration of these knowledge representation formalisms are focused mainly on calculation of probabilities of outcomes of some events. Methodology proposed in this paper is focusing mainly towards the IG of possible actions. This is the crucial difference of the proposed methodology with already existing approaches of integration of ontologies and BNs.

3 USING BAYESIAN NETWORK

It is possible to construct BN (1) manually by knowledge acquisition (in interaction with medical experts), (2) automatically by machine learning algorithms (from available medical data sets), where it is possible to learn network structure and conditional probabilities separately.

The machine learning approach is especially useful in the medical domain where it is very hard to explicitly state medical knowledge, and on the other hand there already are plenty of available medical data sets. With arrival of new patient data the network can be updated and improved. If the environment of the network is changed (e.g. the system is applied in another country), new network can be obtained by learning on new data set.

Figure 1 shows a provisional example of BN that was built manually and that is used in the paper for methodology demonstration purposes (from heart
failure domain). Based on defined conditional probabilities it is possible to calculate probabilities (beliefs) of each outcome of each node in the network (e.g. expecting normal ejection fraction in 70% cases).

In cases when physician is uncertain about the diagnosis she should perform additional diagnostic tests. In that case it would be very useful to know which medical tests are the most appropriate in currently observed patient situation. In other words, it is useful to calculate IG of each observation node for each target node. There are many measures which could be appropriate in this situation; the expected decrease of entropy is a measure which is most commonly used (Jagt 2002).

To calculate the entropy of the target node we use the probabilities of all outcomes of the target node:

$$\text{Entropy}(X) = E(X) = \sum_{o} - p_o \log_2 p_o$$

where $X$ is a target node and $o$ is the outcome of the target node. The maximum value of entropy is 1 (when considering only two possible outcomes: Yes and No) and it is reached when the information about the target node is completely uncertain (when $P(\text{Yes}) = P(\text{No}) = 0.5$). As probability of target node approaches towards ends (0 and 1) the entropy is falls into zero. It is better to have the entropy values as close to zero as possible since that indicates that the answer to the target question is clearer.

A summary measure which takes all possible outcomes of diagnostic test into account is called expected entropy which is calculated as follows:

$$\text{ExpectedEntropy}(X, D) = \sum_{d} p_d E(X \mid D = d),$$

where $X$ is target node, $D$ is observation node, $d$ is a single outcome of node $D$, $p_d$ is probability of occurrence of $d$ outcome, and $E(X \mid D = d)$ is the entropy of the target node $X$ when outcome $d$ has happened. It can be shown that in any BN value of expected entropy cannot be raised by any diagnostic test, only lowered (Jagt 2002).

When the procedure described above is repeated for all observation nodes, a diagram shown on Figure 2 is obtained. The figure indicates that for reaching the final decision whether patient has or has not diastolic heart failure the most informative diagnostic tests is measuring diastolic blood pressure. After physician actually measures diastolic blood pressure beliefs in the network are updated. Accordingly, observation nodes are updated with fresh IG values. Such reasoning procedure in BN is referenced in the literature as “explaining away”.

### 4 STRUCTURED MEDICAL DIAGNOSTIC TESTS

In medical practice some diagnostic measurements are never performed separately (e.g. systolic and diastolic blood pressure). Ontology provides a framework for organizing all possible diagnostic tests into groups as they appear in medical practice. In the ontology grouped diagnostic tests are organized easily by arranging the ontology structure as shown on Figure 3.

![Figure 2: IG of all diagnostic nodes for target concept DIASTOLICHF, for a patient that has not performed a single diagnostic test yet.](image2)

![Figure 3: Diagnostic tests are defined within ontology.](image3)
ECG. Grouped diagnostic tests are not necessarily disjunctive.

4.1 Outcomes of Structured Tests

Difficulty with structured diagnostic tests is that the number of possible outcomes grows extremely fast. Namely, it is equal to the product of number of outcomes for every elementary test in the group. For example, if group blood pressure measurement has only two elementary measurements (systolic and diastolic) where each has three possible outcomes (high, normal, low), there are nine possible outcomes of such test. It is evident that the growth rate of total number of outcomes in the group is of combinatorial nature.

4.2 Information Gain of Structured Medical Test

Formally speaking, to calculate the IG of a group for every possible outcome $g$ of the group $G$ one must calculate (1) the a priori probability of occurrence of observed group outcome $p_g$, and (2) entropy of the target concept when observed outcome $g$ happens $E_g = E(X | G = g)$. Then the expected entropy of the target concept is equal to:

$$\text{ExpectedEntropy}(X, G) = \sum_p p_g E_g$$

where $X$ is observed target node and $G$ is observed grouped diagnostic test.

To find exact value of probability $p_g$ it is possible to construct a dummy node in the BN which would have all nodes from the observed group as parent nodes and conditional probability table defined as truth table which evaluates to YES only in the column of the observed outcome $g$. When network beliefs are updated the belief of outcome YES in that node will be equal to $p_g$. By setting the evidence on the same dummy node to the outcome YES one could read out the a posteriori probability of the observed target node and thus calculate value $E_g$. This procedure should be repeated (1) for every possible outcome, (2) of every possible diagnostic group test, and (3) for every possible target node. With large number of target nodes, large number of grouped diagnostic tests and large number of possible outcomes the procedure becomes extremely computationally demanding. This calls for other potential solutions which would compute in more acceptable time.

One possibility for solving this issue is to use the sampling algorithms. Namely, it is possible to generate arbitrarily large set of samples (artificial set of patients) depending on the properties of the BN and depending on patient evidence that is present and to use it to calculate required probability values. The same procedure applies with grouped tests:

$$p_g = \frac{\text{number of samples with outcome } g}{\text{number of samples}}$$

Table 1 explicates the procedure for computing the expected entropy of the target concept with respect to the grouped diagnostic test.

<table>
<thead>
<tr>
<th>Outcome of grouped test</th>
<th>Probability of outcome $p_g$</th>
<th>Entropy of target concept $E_g$</th>
<th>Contribution to the expected entropy $p_g E_g$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.193</td>
<td>0.986</td>
<td>0.184</td>
</tr>
<tr>
<td>2</td>
<td>0.038</td>
<td>0.750</td>
<td>0.028</td>
</tr>
<tr>
<td>3</td>
<td>0.018</td>
<td>0.601</td>
<td>0.012</td>
</tr>
<tr>
<td>4</td>
<td>0.106</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>5</td>
<td>0.262</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>6</td>
<td>0.064</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>7</td>
<td>0.033</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>8</td>
<td>0.074</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>9</td>
<td>0.213</td>
<td>0.000</td>
<td>0.000</td>
</tr>
</tbody>
</table>

$$\sum_g p_g E_g = 1.0$$

$$\sum_g p_g = 0.231$$

This way the initial set of samples has been divided (unevenly) into nine disjoint subsets. In each subset it is possible to count samples for which the target node was assigned with positive diagnosis. This way a posteriori target probabilities are also calculated from the same sample set:

<table>
<thead>
<tr>
<th>Outcome (g)</th>
<th>Samples</th>
<th>Target Prob.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>19,333</td>
<td>0.193</td>
</tr>
<tr>
<td>2</td>
<td>3,791</td>
<td>0.214</td>
</tr>
<tr>
<td>3</td>
<td>1,779</td>
<td>0.171</td>
</tr>
</tbody>
</table>

Now it is also possible to calculate the a posteriori entropy values and also the final IG value.

5 USAGE EXAMPLE

By starting the decision support services physician finds out probabilities of diseases for the observed patient considering all currently known patient data. An example is shown on Figure 4. When the analysis of IG for all defined grouped diagnostic
tests in the ontology is performed, the physician can find out which test is the most informative one considering observed target nodes. Figure 5 shows the example results of such analysis.

<table>
<thead>
<tr>
<th>Target Node</th>
<th>IG Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diastolic Heart Failure</td>
<td>9%</td>
</tr>
<tr>
<td>Systolic Heart Failure</td>
<td>28%</td>
</tr>
<tr>
<td>Heart Failure Signs</td>
<td>60%</td>
</tr>
<tr>
<td>Heart Failure Symptoms</td>
<td>66%</td>
</tr>
</tbody>
</table>

Figure 5: IG of ontologically organized diagnostic tests considering the target concept “Heart failure signs”.

Medical interpretation of results shown on Figure 5 is following: considering all his currently known data and considering all previously recorded cases of the disease (cases which are inherently encoded within BN) observed patient has 60% possibility of having heart failure signs. If physician wants to be more certain he can perform some additional medical tests. Analysis of IG (again, considering all known patient data and considering all previously recorded cases) indicates that physical examination is the most informative test that can be performed.

Physical examination contains more diagnostic tests one of which is measuring blood pressure, which can be further divided into measuring diastolic and systolic blood pressure (structure defined within the ontology). All such tests (both grouped and elementary) have their own IG value.

A different view of the results is also possible: one can compare the summary impacts of grouped diagnostic tests on defined target concept. Figure 6 demonstrates the comparison of two diagnostic tests: echocardiography and physical examination. This way physician can compare the overall values of IG of all medical tests which helps him to make a decision which medical test (tests) should be performed next.

6 PERFORMANCE

Behaviour of the system in a great deal depends upon some inherent characteristic both of the ontology and the BN. Within this paper measurements are conducted using a single specific BN and a single specific ontology; hence, the analysis is merely a preview of some provisional setting. However, we assume that behaviour of a single problem instance at least to some extent indicates its general behaviour.

The main concern in the performance of the described system is with (1) time required for reasoning and (2) error made in reasoning. Furthermore, it is evident that there is a trade-off between those two parameters.

Figure 7 shows the dependency of error made in reasoning upon number of elementary measurements in the test and the number of samples.

Test: Echocardiography
- Diastolic Heart Failure: 74%
- Systolic Heart Failure: 35%
- Heart Failure Signs: 14%
- Heart Failure Symptoms: 10%

Test: Physical Examination
- Heart Failure Symptoms: 100%
- Heart Failure Signs: 85%
- Systolic Heart Failure: 40%
- Diastolic Heart Failure: 25%

Figure 6: Comprehensive view of IG of available medical diagnostic tests.
standard error made in calculations will be somewhere near 1%.

Figure 8: The appropriate number of samples depends on used number of elementary measurements in the tests and on chosen error rate.

Figure 8 is indicating a minimum number of samples one should use depending on sizes of defined groups in the ontology and on chosen error rate. E.g., if one is satisfied with error rate of 1% and has up to 12 elementary measurements in a group she should use at least 100,000 samples in reasoning phase. Chart depicts such relation for error rates of 2%, 1% and 0.5%.

7 CONCLUSIONS

In this paper we have demonstrated the approach for integration of knowledge from BNs and ontologies in order to calculate the IG of structured medical test. We strongly believe that the approach is sound and can be very useful in practical medical decision support systems.

The main obstacle in the described methodology appeared to be the combinatorial nature of the number of outcomes in grouped diagnostic tests. However, practical experiments indicate that this obstacle in some cases can be to some extent avoided by usage of sampling algorithms in the reasoning phase. The measurements have demonstrated the dependency of error rate and required number of samples. On that basis, and considering some specific system properties such as number of nodes in the network, sizes of grouped diagnostic tests, acceptable time of reasoning, acceptable error rate, and properties of machine that performs reasoning, one can conclude which number of samples should she use in the reasoning phase.

Structure of BNs inherently assumes conditional independency – an assumption that in general case does not stand for medical diagnostic tests. In spite of that, vast majority of decision support systems that make use of BNs ignore this issue. However, one should be fully aware of this drawback when using proposed methodology in practice.

Suggested methodology of integration of BNs and ontologies still calls for more thorough testing of its overall performance and has yet to demonstrate its practical utility in real medical environments. Furthermore, suitability of the approach in some other domains remains to be shown. All above mentioned problems seem to be rather interesting topics for the future work.

REFERENCES