INTRODUCTION

*Haemonchus contortus* is a nematode parasite found in abomasum of sheep and goat, and feeds on blood (Edwards et al., 2016; Wang et al., 2017; Almeida et al., 2018). *H. contortus* infection causes huge economic losses, profitability reduction of the ruminant industry due to weight loss, acute anemia, edema, diarrhea, and lowering in milk and wool production (Emery et al., 2016; Gadahi et al., 2016a). This infection becomes a life threatening disease disturbing sheep population (Besier et al., 2016), because of massive mortality and morbidity rate in affected young animals (Dey et al., 2018). The mortality in lambs and kids in acute cases was ranging from 30% to 50% (Fawzi et al., 2014; Muchiuta et al., 2019). The worm anthelmintic resistance is one of the factors that have supported this economic loss (Borges et al., 2020).
The specificity of carbohydrate to lectins demonstrated that D-mannose, α-D-glucose, and D-N-acetylglucosamin are the major carbohydrate epitopes on *H. contortus* proteins. Therefore, the major part of the carbohydrate moieties on these glycoproteins formed from N-linked oligosaccharides. These indicate that the host immune response against these glycoproteins 55 kDa has ability to inhibit host neutrophils (Anbu and Joshi, 2008).

The current study aimed to present successful glycoprotein fraction for accurate diagnosis of sheep *H. contortus* infection by indirect ELISA.

**MATERIALS AND METHODS**

**Ethical statement**

*H. contortus* adult worms were recovered from slaughtered sheep in the government abattoir according to governmental regulations.

**Parasite**

Live *H. contortus* adult worms of both sexes were collected from abomasum of slaughtered sheep from local abattoirs in Egypt; El-Monieb, El-Bassatin and El-Warrak. The collected worms washed with phosphate buffer saline (0.15 M, pH 7.3) to get rid of debris and stored at -20°C (MAFF, 1986).

**Blood samples**

A total of 274 blood samples were collected from slaughtered sheep at different local abattoirs; El-Monieb, El-Bassatin and El-Warrak in Egypt. The blood samples included; 183 random sheep blood samples. Whereas, the rest 91 blood samples including; 23 positive samples were collected from naturally infected sheep where *H. contortus* adult worms were collected from their abomasum used as true positive sheep sera (gold standard). 13 true negative samples from sheep with no adult worms of *H. contortus* in their abomasum and no other infection of parasites by postmortem examination and fecal analysis were used as uninfected control sheep (gold standard). 19 blood samples were collected from infected sheep with strongyloidiasis. In addition, 9 blood samples were collected from sheep infected with coccidiosis. Other, 9 blood samples from infected animals with fasciolosis as proved by their liver infected with *Fasciola gigantica* and fecal examination. Moreover, 18 blood samples were collected from sheep experimentally infected with *Toxoplasma gondii*; RH isolate from Department of Zoonosis, Veterinary Research Division, National Research Centre. All sera were separately stored at -20°C until use.

**Results**

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PREPARATION OF *H. contortus* WORM CRUDE ANTIGEN

Antigen was prepared by homogenization of adult worms in phosphate buffer saline at 4°C using a tissue-glass homogenizer, then sonication three times for 20 s each time at 100 m Amp by 150 ultra-sonication (Sanyo Gallenkamp PLC, UK), then centrifugation for 30 min at 12 000 rpm and 4 °C, then the supernatant was collected and kept at −20 °C (Prasad et al., 2008). The protein content of the collected supernatant (antigen) measured by Lowry method (Lowry et al., 1951).

PURIFICATION OF GLYCOPROTEINS BY LECTIN AFFINITY CHROMATOGRAPHY

The *Concanavalin ensiformis* (Con A) column obtained from Sigma Chem. Co. St. Louis was used for purification of glycoprotein antigens as previously described by Fukuda and Kobata (1993). The crude *H. contortus* antigen was applied to the column and the bound fraction was eluted with four types of sugars separately; glucose, N-acetylglucosamine, galactose and lactose (Gluc- GlucNAc- Gal- and Lac). The protein content of fractions was checked by Lowry method (Lowry et al., 1951).

CHARACTERIZATION OF ISOLATED FRACTIONS

SDS-PAGE

Glycoprotein fractions, crude *H. contortus* antigen and molecular weight protein standards (Sigma, USA) were separately, electrophoresed on 10% gel of SDS- PAGE under reducing condition according to Laemmli (1970). After separation, the gel stained with silver stain according to Wray et al. (1981). The bands were analyzed and photographed by Molecular Imager Gel Doc™XR + with Image Lab Software (Bio-Rad).

WESTERN BLOT

Immunoreactive bands in four isolated fractions and crude extract of *H. contortus* were identified by pooled positive sera of sheep naturally infected with haemonchosis in western blot. The four fractions, crude antigen and prestained protein Ladder (Vivantis Technologies) were blotted onto nitrocellulose membrane as described by Towbin et al. (1979) in a blotting system. The non-specific immunoreactive bands in the most potent diagnostic fractions were also identified in immunoblot using sheep serum samples infected with other parasitic infections as fascioliosis, toxoplasmosis, strongyloidiasis and coccidiosis, anti-sheep IgG horseradish peroxidase conjugate and 4-chloro-1-naphthol (Sigma) were used. Finally, the membrane was analyzed and photographed by Molecular Imager Gel Doc™XR + with Image Lab Software (Bio-Rad).

ELISA

Indirect-ELISA was used to evaluate the diagnostic potency of the four isolated fractions in detecting anti-*H. contortus* IgG in sheep sera naturally infected with haemonchosis. The fraction with highest diagnostic potency was used for diagnosis of haemonchosis in random sheep samples using indirect ELISA. The concentration of conjugate, antigen and tested sera were determined by checkerboard titrations. The optical densities (OD) were read at 450 nm with ELISA reader (BIO-TEK, INC., ELx, 800UV). The cut off value was calculated by mean OD values of negative control ± 3SD (Gowda, 2016).

ESTIMATION OF SENSITIVITY, SPECIFICITY AND PREDICTIVE VALUES OF THE MOST POTENT DIAGNOSTIC FRACTION

The cross reactivity with other parasitic infections than haemonchosis as strongyloidiasis (*Strongyloides papillosus*), fascioliosis (*Fasciola gigantica*), toxoplasmosis (*T. gondii*) and coccidiosis (*Eimeria* spp.) was estimated in the presence of positive sheep sera with haemonchosis (*H. contortus*) and negative sera using indirect ELISA. The sensitivity, specificity and predictive values were estimated as following formula (Parikh et al., 2008; Deo et al., 2019).

\[
\text{Sensitivity} = \frac{tp}{tp + fn} \\
\text{Specificity} = \frac{tn}{tp + fp + tn + fn} \\
\text{Positive Predictive Value (PPV)} = \frac{tp}{tp + fp} \\
\text{Negative Predictive Value (NPV)} = \frac{tn}{tn + fn} \\
\text{Diagnostic Efficacy=} \frac{Ntn + Ntp}{(Ntp + Nfn) + (Ntp + Nfp + Ntn + Nfn)}
\]

Whereas, \(N\) (number of samples), \(tp\) (true positive), \(fp\) (false positive), \(tn\) (true negative) and \(fn\) (false negative).

STATISTICAL ANALYSIS

The optical densities data were expressed as arithmetic mean ± standard deviation. The apparent prevalence parameter was analyzed using Chi square test by statistical computer package for social science (SPSS) version 15.

RESULTS

LECTIN AFFINITY CHROMATOGRAPHY PROFILE OF *H. contortus* ADULT WORM ANTIGEN

Purification of *H. contortus* adult worms antigen by con A-affinity column chromatography resulted in four fractions; D- Glucose, D- Galactose, N-Acetylgalactosamine and Lactose fractions. The protein content of Gluc fraction was 78.3µg/ml, Gal-fraction was 57.9µg/ml, GlucNAc– fraction 62.5µg/ml and Lac fraction 60.3µg/ml, whereas, protein content of crude extract was 330.4 µg/ml.

ELECTROPHORETIC PROFILE OF *H. contortus* FRACTIONS

The electrophoresis profile of crude antigen showed 9 bands with a wide range of molecular weights, but the
bands 100, 83, 70, 66, 56, 36, 31, 27 and 24 kDa seem to be the predominant bands as shown in Figure 1 Lane A. GlucNAc fraction separated into three bands with molecular weight 83, 66 and 24 kDa (Lane B). Gal fraction resolved into four bands of 87, 70, 36 and 27 kDa (Lane C). Gluc fraction displayed three bands with molecular weight 83, 66 and 24 kDa (Lane D). Lactose fraction was represented by two bands at molecular weight 87 and 26 kDa (Lane E) (Figure 1).

**Figure 1:** Electrophoretic profile of crude antigen of *H. contortus* adult worm (Lane A) and glycoproteins fractions; GlucNAc (Lane B), Gal (Lane C), Gluc (Lane D) and Lac (Lane E). Molecular weight protein standard (Lane St).

**Western Blot**
The specific 24 kDa immunogenic reactive band in both Gluc and GlucNAc fractions reacted strongly with positive sheep sera naturally infected with *H. contortus*. Whereas, Gal and Lac fractions were very weakly reactive. The crude antigen was reactive with anti-*H. contortus* at four immunogenic bands 100, 83, 36, and 24 KDa (Figure 2).

**Diagnostic Potentials of *H. contortus* Fractions**
The diagnostic profile of the fractions showed the highest diagnostic potentials of Gluc fraction than other fractions using highly diluted antibodies 1:65536 (Figure 3).

**Figure 2:** Immunogenic bands of *H. contortus* antigens identified by specific naturally infected sheep sera by Western Blot assay. Crude *H. contortus* antigen (Lane A), GlucNAc (Lane B), Gal (Lane C), Gluc (Lane D) and Lac (Lane E). Pre stained molecular weight Marker (Lane MW.).

**Figure 3:** Diagnostic profile of four fractions compared to crude antigen in natural sheep haemonchosis. The difference is significant p ≤ 0.05.
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The success of Gluc fraction in detection of specific haemonchus IgG in sheep was assessed by indirect ELISA using different antibodies with other parasitic infections. The fraction recorded 97.05% specificity (Figure 4). Whereas, the fraction recorded 5.8% cross reactivity (false positive) with sheep serum infected with strongyloidiasis (n=19, one false positive for haemonchosis) and 11% with fasciolosis IgG (n=9, one false positive for haemonchosis). No cross reactivity was recorded with uninfected sheep sera (negative sera for *H. contortus*), and no cross reactivity with antibodies of toxoplasmosis and coccidiosis, (Figure 4). Serum samples of sheep infected with *H. contortus*, as proved by post mortem examination, reacted positively with Gluc fraction recording 100% sensitivity (Figure 4).

The fraction recorded Positive Predictive Value (PPV) 92%, Negative Predictive Value (NPV) 100 % and diagnostic efficacy 97.8 %. The current results of sensitivity and specificity were confirmed by western blot (Figure 5).

**Sheep Haemonchosis Infection Percentage by Indirect ELISA Using Gluc Fraction**
The assay recorded 56.83% infection percentage in the examined sample. OD values of the examined serum samples ranged from 0.045 to 0.795 (Figure 6).

**DISCUSSION**

In the current study, four glycoprotein fractions were isolated from whole worm crude extract. Electrophoretic profile of the four fractions showed that both Gluc and GlucNAc fractions have common three bands with molecular weight 83, 66 and 24 KDa. While Gal fraction resolved into four bands 87, 70, 36 and 27 KDa, and Lac fraction was represented by two bands at molecular weight 87 KDa and 26 KDa. Whereas, crude extract antigen separated into many bands at molecular weight range from 100 kDa to 24 KDa. These bands of crude extract were found in the range (14 to 216 kDa) previously described by Derbala and Abd El-Rahman (2001). Whereas, Anbu and Joshi (2008) isolated 55 kDa secretory glycoprotein (gp55) from E/S products of adult *H. contortus* by using Con a Sepharose column. While, Ashman et al. (1995) purified 70-90 kDa larval surface glycoprotein of *H. contortus* by size exclusion chromatography. Moreover, the galectin 11 and 14 ligands were isolated from *H. contortus* by the galectin affinity column (Sakhivie et al., 2018). Whereas, Naqvi et al. (2020b) observed that rHc-Cs which purified as Histidine-tagged fusion protein, resolved into single band at 38 kDa, while native CS protein resolved at 20 kDa.

**Figure 4:** Specificity, sensitivity and predictive values of Gluc fraction by indirect ELISA.

**Figure 5:** Immunoblotting of isolated Gluc fraction treated with; positive serum with haemonchosis (lane1), (strongyloidiasis (Lane 2), fasciolosis (Lane 3), toxoplasmosis (Lane 4), coccidiosis (Lane 5) and negative control (Lane 6). Pre stained molecular weight Marker (Lane Mw).

**Figure 6:** Diagnostic potency of Gluc fraction in randomly collected sheep sera.
on 12% SDS-PAGE. The difference in molecular weight of glycoproteins bands may be due to the difference in origin of crude extract from which glycoproteins isolated, or antigen preparation, eluting buffers and type of column used in purification process.

In the current study, 24 kDa is immunoreactive band of that found in both Gluc and Gluc-NAc fractions which showed highest diagnostic potentials. This common band is mainly responsible for the diagnostic activities in both fractions. The difference in diagnostic properties of Gluc and Gluc-NAc fractions may be attributed to quantities of this band in both fractions. The band 24 kDa was previously isolated from *H. contortus* excretory-secretory products of adult worm (HeES-24) (Gadahi et al., 2016). And diagnostic 26-kD band was isolated from adult *H. contortus* by anion exchange chromatography and proved potency in early and late experimental sheep haemonchosis (Gomez-Munoz et al., 2000; Arab et al. (2013) proved the potency of 26 KD which isolated from adult *H. contortus* by gel filtration chromatography using sephadex G 100 in the diagnosis of sheep haemonchosis.

In the present study Gluc fraction has been successfully detected specific antibodies of *H. contortus* with specificity 97.05% and sensitivity 100% using indirect ELISA. The results of specificity were confirmed by western blot. Antigen specificity is essential in accurate sero-diagnosis of diseases. This fact can be applied with all pathogens particularly parasites. Where cross reaction is a common trait among helminthes (Abdel-Rahman and Abdel-Meaged, 2000; Hassan et al., 2012; Shaapan et al., 2015) and consequently, isolation of specific antigen is a challenge.

In previous study, Naqvi et al. (2019) showed that no cross reactivity with the purified rHc-ICAS99 antigen from the excretory and secretory products against infected sera with *T. spiralis*, *F. hepatica* and *T. gondii*. Moreover, our results agree with Naqvi et al. (2020b) who reported that western blotting results showed that rHc-TpMy was detected all positive sera of haemonchosis while no antibody detection was observed against negative sera. In addition, Naqvi et al. (2019) reported that combined use of western blot and indirect ELISA analysis are potential diagnostic tool for diagnosis of *H. contortus* infection in goat. In contrast to this, Kandil et al. (2017a) showed that there were cross reactivity with other cestodes,*Moniezia expansa* and *F. hepatica* against glutathione-S-transferase (GST) and recombinant protein (rhp 26/23) antigens based on both indirect ELISA and immunoblot assays. Moreover, partially purified crude *H. contortus* antigen by gel filtration using Sephadex G-100, showed cross reactivity with other cestodes and *F. hepatica* in both indirect ELISA and immunoblot (Kandil et al., 2015). In addition, Kandil et al. (2017a) demonstrated that the recombinant *H. contortus* glutathione-S-transferase was showed high false positive with negative sheep sera and scored highest prevalence (90.8%) and sensitivity (90%) by indirect ELISA. Therefore, Wartini et al. (2017) explained that False positive/negative results are unwanted and may affect the efficacy of assays so, they can performed the optimization of antigen/antibodies concentration to improve the diagnostic potential of assay.

In current study Gluc-glycoprotein fraction record sensitivity 100% and specificity 97.05% which is higher than somatic antigen which recorded specificity 67.18% by indirect ELISA in previous study (Gowda, 2016). And it may be similar to rHc-TpMy-which showed 90% sensitivity and 100% specificity (Naqvi et al., 2020b). While, Song et al. (2018) consider that the sensitivity and specificity of assay affected by the type of antigens used. Whereas, Gomez et al. (1995, 2000) and Arab et al. (2013) suggested that the 26 kDa antigen of adult *H. contortus* has been seem specific for the diagnosis of *H. contortus* infection. On the other hand, Mir et al. (2008) showed that adult crude antigen scrod 72.22 % and 76.81 % sensitivity and specificity respectively by indirect ELISA. While, Sultan et al. (2012) reported that the sensitivity of crude adult worm antigen was 87.5% and specificity 75 %. Moreover, previous study Schallig et al. (1995) showed that crude somatic antigens based on ELISA recorded sensitivity 89.2 % and specificity 82.7 % for diagnosis haemonchosis in sheep.

The infection percentage of haemonchosis in the current study is 56.83%, Positive Predictive Value 92%, Negative Predictive Value was 100 % and 97.8 % diagnostic efficacy using indirect ELISA. These results indicate that this test indirect ELISA based on new Gluc-glycoprotein fraction is doing as good as “gold standard”. And other fractions recorded in previous studies, different infection percentages were recorded based on the used antigens and assays. 72.22 % sero-prevalence was observed with crude *H. contortus* somatic antigen (Mir et al., 2008). 58.66 % serum samples screened using somatic antigen (Gowda, 2016). Whereas, Hassan et al. (2019) reported that the sero-prevalence of haemonchosis among sheep was higher than goats and represented 64.48% and 41.91% respectively. Additional factor that is responsible for different infection percentages is body condition where, Brik et al. (2019) reported maximum prevalence 80.83% in sheep with poor body, (34.26%) in moderate body and 6.88% in sheep with good body. The difference could be also attributed to examine sample size.

**CONCLUSION**

Gluc fraction isolated from *H. contortus* adult worm antigen by Con A lectin column chromatography with immunoreactive band 24kDa was successfully utilized in
the diagnosis of sheep haemonchosis with 100% sensitivity, 97.05% specificity, Positive Predictive Value 92%, and Negative Predictive Value 100 % with 97.8 % diagnostic efficacy using indirect ELISA. The immunogenic band of 24 kDa is mainly responsible for its potency. Further purification and characterization processes are needed to increase the specificity of the fraction and its diagnostic potency.

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AUTHOR’S CONTRIBUTIONS

The authors participated in planning and study design. TNI collected *H. contortus* adult worms from abomasum of slaughtered sheep and blood samples. TNI prepared crude antigen, performed lectin chromatography, SDS-PAGE and western blot assays. TNI and A-REH shared in ELISA. TNI and A-REH shared in analyzed data and writing the manuscript.

CONFLICT OF INTEREST

The authors have declared no conflict of interest.

REFERENCES


