INTRODUCTION

Research focused on decreasing enteric methane production from ruminants has been reported in the last decade, with various mitigation strategies available to the livestock fields (Knapp et al., 2014). Enteric methane production from ruminal fermentation of feeds accounts for approximately 30% of the total anthropogenic methane (FAO, 2006). Methane production from ruminants is a serious contributor to global warming as well as an economic loss to the farmer with the loss of 2% to 15% of the gross energy intake through their unique digestive process (Johnson and Johnson, 1995; Ellis et al., 2007). Rumen nutritionists have found bio-active materials for reducing methane production without an adverse effect on ruminal fermentation. Natural sources including plant extracts and essential oils and marine algae have been studied and screened as additives for ruminants feeds (Dubois et al., 2013; Durmic et al., 2014), but only a few substances have been shown to have an effect on ruminal methane production. Plant ex tracts such as condensed tannin, saponins and essential oils modify the ruminal ecosystem by suppressing ruminal protozoa, and inhibiting bacteria and archaea (Tan et al., 2011; Patra and Yu, 2012). According to above studies, research on the symbiosis of protozoa with methanogens in the rumen is well reported, and the inhibition of protozoa has been shown to be an effective approach to decrease ruminal methane production. However, there are limited studies about the effect of medicinal herb extracts (MHEs) on ruminal methanogenesis. This study was aimed to evaluate the effect of MHEs on
the in vitro ruminal methane production, fermentation characteristics and how some plants secondary metabolites might act as inhibitors of protozoa closely related with methanogens in the rumen.

MATERIALS AND METHODS

Preparation of medical herb extracts

MHEs were obtained from Plant Extract Bank at the Korea Research Institute of Bioscience and Biotechnology (Daejeon, Korea). Plants were collected from fields in Korea (Table 1). Each plant was cut into small pieces and dried naturally under shade. The dried plants (100 g) were extracted with 99.9% methyl alcohol (1 L) using ultrasonic cleaner (Branson Ultrasonics corporation, Danbury, CT, USA) at room temperature for 3 days. After extraction, the solutions were filtered and the solvents were evaporated under vacuum. Stock solutions (20 mg/mL) of the extract were dissolved in dimethyl sulfoxide (Sigma-Aldrich Chemical Co., St. Louis, MO, USA) and diluted using culture media immediately before experiments.

Ruminal inoculum and in vitro incubation

A fistulated Hanwoo cow (three years old) of 450 kg body weight was used as the donor of rumen fluid. Timothy and commercial concentrate in the ratio of 60:40 were fed at 3% of body weight twice a day (09:00 and 17:00). Water and mineral-vitamin block were allowed ad libitum. The rumen fluid was collected from the fistulatedHanwoo before morning feeding. Rumen liquor was filtered through four layers of cheesecloth before mixing with buffer and was maintained at 39°C. The 15 mL of rumen fluid-buffer mixture, comprising McDougall buffer (McDougall, 1948) and rumen liquor in the ratio of 2 to 1, was dispensed anaerobically into 50 mL serum bottles containing 0.3 g of timothy substrate and MHEs (5% of substrate). The serum bottles were filled with O2-free N2 gas capped with a rubber stopper and held in a shaking incubator (Jeio Tech, SI-900R, Korea; 120×rpm) at 39°C for 12 h, 24 h, and 48 h. The incubated samples taken from each fermentation period were centrifuged at 3,000×rpm for 3 min to remove feed particles, and the supernatants were re-centrifuged at 14,000×rpm for 3 min to settle the pellets down. After that, sodium phosphate buffer (pH 6.5) was added to these precipitates and vortexed. Growth rates of total microbes were estimated as optical density values using spectrophotometer (Model 680, Bio-Rad Laboratories, USA). The culture was subsampled for the analysis of pH (Mettler-Toledo, MP230, Greifensee, Switzerland), volatile fatty acid (VFA) concentration and genomic DNA extraction. The VFA analysis was performed with a high performance liquid chromatography (Agilent-1200, Waldbronn, Germany) equipped with column (300 mm×7.8 mm I.d. MetaCarb 87H, Varian, Palo Alto, CA, USA). In vitro dry matter disappearance rate was estimated by the modified method of nylon bag digestion process. Briefly, after incubation, the nylon bag with substrate was washed twice in a water-bath equipped with Heidolhs Rotamax 120 (Heidolph Instrument, Nuremberg, Germany) at 100×rpm for 30 min. Washed nylon bags were then dried to a constant weight at 60°C. Dry matter disappearance was determined by weight difference before and after incubation in the serum bottle.

Microbial growth rate

Incubated samples taken from each fermentation period were centrifuged at 3,000×rpm for 3 min to remove feed particles, and the supernatants were re-centrifuged at 14,000×rpm for 3 min to settle the pellets down. After that, sodium phosphate buffer (pH 6.5) was added to these precipitates and vortexed. Growth rates of total microbes were estimated as optical density values using spectrophotometer (Model 680, Bio-Rad Laboratories, USA). The culture was subsampled for the analysis of pH (Mettler-Toledo, MP230, Greifensee, Switzerland), volatile fatty acid (VFA) concentration and genomic DNA extraction. The VFA analysis was performed with a high performance liquid chromatography (Agilent-1200, Waldbronn, Germany) equipped with column (300 mm×7.8 mm I.d. MetaCarb 87H, Varian, Palo Alto, CA, USA). In vitro dry matter disappearance rate was estimated by the modified method of nylon bag digestion process. Briefly, after incubation, the nylon bag with substrate was washed twice in a water-bath equipped with Heidolhs Rotamax 120 (Heidolph Instrument, Nuremberg, Germany) at 100×rpm for 30 min. Washed nylon bags were then dried to a constant weight at 60°C. Dry matter disappearance was determined by weight difference before and after incubation in the serum bottle.

Gas production measurement, analysis of gas and ruminal fermentation profiles

At the end of incubation, total gas production was measured according to the assay outlined by Theodorou et al. (1994). A detachable pressure transducer and a digital readout voltmeter (Laurel Electronics, Inc., Costa Mesa, CA, USA) were used to measure the headspace gas pressure of fermenting cultures. For the total gas production measurement, the transducer was modified in a way that it could be linked to the inlet of a disposable Luer-lock threeway stopcock (Theodorou et al., 1994). Gas pressure in the headspace was read from the display unit after insertion of the hypodermic syringe needle through the butyl rubber stopper above the culture medium. The headspace gas in the serum bottle was collected for analyzing methane, carbon dioxide and hydrogen by gas chromatography (Agilent Technologies HP 5890, Santa Clara, CA, USA) conducted using a TCD detector with a Column Carboboxen 1006PLOT capillary column 30 m×0.53 mm (Supelco, Bellefonte, PA, USA). The culture was subsampled for the analysis of pH (Mettler-Toledo, MP230, Greifensee, Switzerland), volatile fatty acid (VFA) concentration and genomic DNA extraction. The VFA analysis was performed with a high performance liquid chromatography (Agilent-1200, Waldbronn, Germany) equipped with column (300 mm×7.8 mm I.d. MetaCarb 87H, Varian, Palo Alto, CA, USA). In vitro dry matter disappearance rate was estimated by the modified method of nylon bag digestion process. Briefly, after incubation, the nylon bag with substrate was washed twice in a water-bath equipped with Heidolhs Rotamax 120 (Heidolph Instrument, Nuremberg, Germany) at 100×rpm for 30 min. Washed nylon bags were then dried to a constant weight at 60°C. Dry matter disappearance was determined by weight difference before and after incubation in the serum bottle.

Table 1. Technical information regarding medicinal herb extracts used in this study

| Stock No. | Scientific name         | Family     | Sampling period |
|-----------|-------------------------|------------|----------------|
| 004-034   | Veratum patulum         | Liliaceae  | May 11, 2001   |
| 012-017   | Iris ensata var. spontanea | Iridaceae  | September 9, 2001 |
| 019-088   | Arisaema ringens        | Araceae    | October 11, 2002 |
| 028-080   | Carduus crispus         | Compositae | May 18, 2005   |
| 036-088   | Pueraria thunbergiana    | Leguminosae| August 6, 2009 |

Medical herb extracts were obtained from Plant Extract Bank (PEB) at Korea Research Institute of Bioscience and Biotechnology (Daejeon, Korea).
Quantitative polymerase chain reaction assays

DNA extraction: A high-speed reciprocal shaker which retains samples in screw-capped tubes containing silica beads was used for DNA extraction. Total nucleic acid was extracted from the incubated rumen samples by using the modified bead-beating protocol with the Soil kit (Macherey-nagel, Düren, Germany). Briefly 1.0 mL aliquot was taken from the incubated culture solution and was centrifuged at 3,000×rpm. Nucleic acid concentrations were measured by using a NanoDrop Spectrophotometer (Thermo Scientific, Wilmington, DE, USA).

Polymerase chain reaction primers: The polymerase chain reaction (PCR) primer sets used in this study for amplification of ciliate protozoa (Forward primer: GAG CTA ATA CAT GCT AAG GC, Reverse primer: CCC TCA CTA CAA TCG AGA TTT AAG G), Fibrobacter succinogenes (Forward primer: GTT CGG AAT TAC TGG GCC TAA A, Reverse primer: CGG CTG CCC CTG AAC TAT C), Ruminococcus ldlavaciens (Forward primer: CGA ACG GAG ATA ATT TGA GTT TAC TTA GG, Reverse primer: CGG TCT CTG TAT GTT ATG AGG TAT TAC C) and Ruminococcus albus (Forward primer: CCC TAA AAG CAG TCT TAG TTC G, Reverse primer: CCT CCT TGC TAC TCG CAT A) were the same as those referenced by Skillman et al. (2006), Denman and McSweeney (2006) and Koike and Kobayashi (2001), respectively. Total bacteria primer set (Forward-CGGCAACGAGCG CAACCC, Reverse-CCATTGTAGCACGTGTGTAGCC) was used as the internal standard (Denman and McSweeney, 2006).

Real-time PCR: Real-time PCR (RT-PCR) assays for enumeration of microbes were performed according to the methods described by Denman and McSweeney (2006) and Denman et al. (2007) on a real-time PCR Machine (CFX96 Real-Time system, BIO RAD, Hercules, CA, USA) using the SYBR Green Supermix (QPK-201, Toyobo Co., LTD., Tokyo, Japan). The values of cycle threshold (Ct) after PCR reactions were used to determine fold change (number of fold difference) of different microbial population relative to control without additives. Abundance of these microbes was expressed by the equation:

\[
\text{Relative quantification} = 2^{- \Delta \text{Ct(Target)} - \Delta \text{Ct(Control)}}
\]

Where, Ct represents threshold cycle. All RT-PCR reaction mixtures (final volume of 20 μL) contained forward and reverse primers, the SYBR Green Supermix and DNA template. A negative control without the template DNA was used in every RT-PCR assay for each primer. The RT-PCR amplification of the target DNA, included the annealing and the extension temperature, was performed as previously described (Koike and Kobayashi, 2001; Denman and McSweeney, 2006; Skillman et al., 2006).

Statistical analysis

Data were analyzed using the general linear model procedure of the Statistical Analysis System Institute, Inc. (SAS Institute, 1996). The effects of MHEs on pH, VFA, dry matter degradability, total gas production, gas profiles and microbial growth were compared to the controls and significant differences between treatment means were examined using Duncan’s multiple comparison tests. A p<0.05 was considered to be statistically significant.

RESULTS AND DISCUSSION

The in vitro ruminal fermentation characteristics are shown in Table 2. The pH of culture solution did not show a significant difference between control and treatments (p>0.05). Dry matter degradability was not significantly different, but numerically higher compared with the control. The total volatile fatty acid (TVFA) concentrations were decreased or increased by addition of MHEs at 12 h incubation, while they were significantly increased (p<0.05) compared to the control at 24 h incubation except Veratrum patulum. The acetate:propionate (C2:C3) ratio was lowered by the addition of all MHEs compared to the control at 24 h incubation. The effects of MHEs on total gas production and gas profiles are presented in Table 3. Total gas production was higher than that of the control. There was significant difference (p<0.05) at 48 h incubation. MHEs decreased the in vitro methane production at 12 h and 24 h incubation. In particular, Iris ensata var. spontanea and Arisaema ringens extracts significantly reduced (p<0.05) the in vitro methane production by 39.6% after 12 h incubation and 43.9% after 24 h incubation, respectively. Carbon dioxide production was not significantly different. Unfortunately, hydrogen was not detected in this in vitro study. According to the study of Janssen (2010), it is difficult to accumulate hydrogen in the ruminal ecosystem. The normally low concentration is accounted for by the utilization of hydrogen by ruminal methanogens. Carroll and Hungate (1955) also reported that hydrogen was metabolized quickly, with stoichiometric methane production when added to in vitro ruminal fermentation. Some researchers reported the rate of methane production and the concentrations of dissolved hydrogen instead of hydrogen as an intermediate in rumen fermentation (Hungate, 1967; Barry et al., 1977; Hillman et al., 1985). In addition, Vogels et al. (1980) reported that hydrogen production is closely related with anaerobic ciliates. The growth of the rumen microbes as affected by all treatments is presented in Table 4. The result of microbial growth rate was not significant difference between the control and
The effect of MHEs on ruminal microbe diversities at 24 h incubation is shown in Figure 1. The protozoa population in all added MHEs was shown to decrease dramatically (p<0.05) as compared with the control (Figure 1a). From the study of Eom et al. (2013), plants of *Iridaceae* in Korea contain tannin and various kinds of terpenoids. According to previous studies, tannins and saponins inhibited ruminal protozoa diversity, and the reduced methane production might be related with a decreased protozoa population (Makkar et al., 1995; Hristov et al., 2003). This finding agreed with the present results that *Iris ensata var. spontanea* extract led to reduce ruminal methane production by 39.6% and 34.8% at 12 h and 24 h incubations, respectively.

### Table 2. Effects of medicinal herb extracts on ruminal fermentation characteristics according to incubation time

| Item            | Control | T1   | T2   | T3   | T4   | T5   | SEM  |
|-----------------|---------|------|------|------|------|------|------|
| DMD (%)         | 27.64   | 27.54| 25.00| 27.58| 26.48| 27.72| 1.70 |
| pH              | 6.61    | 6.62 | 6.63 | 6.63 | 6.65 | 6.62 | 0.03 |
| tVFA (mM)       | 40.56   | 45.42| 38.87| 45.11| 39.87| 47.04| 4.98 |
| Acetate (mM)    | 31.77   | 33.10| 31.36| 33.17| 33.08| 33.45| 1.55 |
| Propionate (mM) | 16.32   | 19.28| 16.87| 16.68| 19.98| 20.97| 4.13 |

SEM: standard error of the mean; DMD, dry matter degradability; tVFA, total volatile fatty acid.

* Control, no addition; T1, *Veratrum patulum*; T2, *Iris ensata var. spontanea*; T3, *Arisaema ringens*; T4, *Carduus crispus*; T5, *Pueraria thunbergiana*.

### Table 3. Effects of medicinal herb extracts on gas production and gas profiles according to incubation time

| Item (mL/g DM) | Control | T1   | T2   | T3   | T4   | T5   | SEM  |
|----------------|---------|------|------|------|------|------|------|
| Gas            | 187.63  | 193.23| 172.16| 177.13| 171.05| 176.39| 3.04 |
| CH₄            | 5.46    | 5.06  | 3.30  | 3.53  | 4.38  | 4.61  | 0.92 |
| CO₂            | 64.87   | 75.26 | 61.19 | 66.90 | 61.20 | 71.74 | 8.53 |

SEM: standard error of the mean.

* Control, no addition; T1, *Veratrum patulum*; T2, *Iris ensata var. spontanea*; T3, *Arisaema ringens*; T4, *Carduus crispus*; T5, *Pueraria thunbergiana*.

*abcd Means in the same row with different superscripts differ significantly (p<0.05).*

treatments. The effect of MHEs on ruminal microbe diversities at 24 h incubation is shown in Figure 1. The protozoa population in all added MHEs was shown to decrease dramatically (p<0.05) as compared with the control (Figure 1a). From the study of Eom et al. (2013), plants of *Iridaceae* in Korea contain tannin and various kinds of terpenoids. According to previous studies, tannins and saponins inhibited ruminal protozoa diversity, and the reduced methane production might be related with a decreased protozoa population (Makkar et al., 1995; Hristov et al., 2003). This finding agreed with the present results that *Iris ensata var. spontanea* extract led to reduce ruminal methane production by 39.6% and 34.8% at 12 h and 24 h incubations, respectively. Among MHEs treatments,
Arisaema ringens and Pueraria thunbergiana, and Carduus crispus used in this study are known to have saponin and flavonoid properties (Kim and Kim, 2003; Lee et al., 2009; Kim et al., 2012). The addition of Arisaema ringens, Pueraria thunbergiana, and Carduus crispus led to decrease the protozoa population in this study. Patra and Saxena (2010) showed that saponins might decrease the ruminal methanogenesis by inhibiting rumen protozoa or the numbers and activity of methanogens with high dosage of saponins although the direct effect of saponins on methanogens has not been reported clearly. Oskoueian et al. (2013) reported that flavonoids inhibited the total protozoa and total methanogens, and decreased ruminal methane production. In addition, the dry matter degradability and other ruminal fermentation characteristics were not affected by flavonoids. Furthermore, the diversities of Fibrobacter succinogenes, Ruminococcus flavefaciens, and Ruminococcus albus considered representative cellulolytic bacterial species in the rumen are shown in Figure 1 (b, c, and d). The F. succinogenes community under MHEs except for Carduus crispus was shown to increase, while the R. flavefaciens and R. albus diversities under all MHEs decreased as compared with the control. In particular, F. succinogenes has been known as an important species for fiber digestion in the rumen. Minato and Suto (1978) showed that F. succinogenes had the highest cellulose attachment ability among ruminal bacteria in vitro. R. flavefaciens and R. albus are known as promising bacteria to produce succinic acid and hydrogen, respectively.

Table 4. Effects of medicinal herb extracts on growth rate of ruminal microbes according to incubation time

| Incubation time (h) | Control$^1$ | T1 | T2 | T3 | T4 | T5 | SEM |
|---------------------|-------------|----|----|----|----|----|-----|
| 12                  | 0.45$^{abc}$| 0.42$^c$| 0.43$^b$| 0.50$^a$| 0.51$^a$| 0.52$^a$| 0.04 |
| 24                  | 0.40        | 0.40| 0.39| 0.40| 0.37| 0.39| 0.03 |
| 48                  | 0.42$^{ab}$| 0.44$^a$| 0.45$^b$| 0.44$^a$| 0.38$^b$| 0.48$^a$| 0.04 |

SEM, standard error of the mean.

$^1$ Control, no addition; T1, Veratrum patulum; T2, Iris ensata var. spontanea; T3, Arisaema ringens; T4, Carduus crispus; T5, Pueraria thunbergiana.

$^a$$^b$$^c$ Means in the same row with different superscripts differ significantly ($p<0.05$).
This finding agreed that the decreased \textit{R. flavefaciens} and \textit{R. albus} populations might be related with the increased propionate and undetected hydrogen production in this study. Moreover, hydrogen produced during enteric fermentation in ruminants is the critical concern to the microbial ecosystem and the regulation of hydrogen is the key to decrease methane production from ruminal fermentation (Hungate, 1967). In addition, the formation of propionate from succinate would affect the lower availability of hydrogen for the ruminal methanogenesis (Mitsumori and Sun, 2008).

In conclusion, this study of screening a variety of native plant extracts including MHEs may provide useful information on the potential candidates for ruminal additives in further studies. The addition of MHEs used in this study had no significant negative effects on the ruminal fermentation characteristics. The \textit{in vitro} ruminal methane production was reduced by addition of MHEs, and these MHEs dramatically suppressed the population of ruminal protozoa. For most of the ruminal additives including plants and plant secondary metabolites, their effects also depend on the diets as well. However, comprehensive \textit{in vivo} studies with ruminants need to be conducted to evaluate the sustainability of MHEs supplementation to reduce the ruminal methane production and methanogenesis without adverse effects on the animal.

CONFLICT OF INTEREST

We certify that there is no conflict of interest with any financial organization regarding the material discussed in the manuscript.

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