Bioprocessing of wheat straw into nutritionally rich and digested cattle feed

Bhuvnesh Shrivastava¹, Kavish Kumar Jain¹, Anup Kalra² & Ramesh Chander Kuhad¹

¹Lignocellulose Biotechnology Laboratory, Department of Microbiology, University of Delhi South Campus, New Delhi 110021, India, ²Ayurvet Ltd., Sagar Plaza, District Centre, Laxmi Nagar, Delhi 110092, India.

Wheat straw was fermented by Crinipellis sp. RCK-1, a lignin degrading fungus, under solid state fermentation conditions. The fungus degraded 18.38% lignin at the expense of 10.37% cellulose within 9 days. However, when wheat straw fermented for different duration was evaluated in vitro, the 5 day fungal fermented wheat straw called here “Biotech Feed” was found to possess 36.74% organic matter digestibility (OMD) and 5.38 (MJ/Kg Dry matter) metabolizable energy (ME). The Biotech Feed was also observed to be significantly enriched with essential amino acids and fungal protein by fungal fermentation, eventually increasing its nutritional value. The Biotech Feed upon in vitro analysis showed potential to replace 50% of concentrate, which was experimentally evaluated by adding Biotech Feed based diets to feed intake of an average 27 kg BW per day, growing and finishing beef calves. When animals were fed with Biotech Feed for 55 days, it was observed that the calves fed on Biotech Feed gained higher than the control fed on concentrate mixture. Further, the calves fed on Biotech Feed based diets exhibited significantly higher (p<0.05) dry matter intake (DMI: 3.74 Kg/d), dry matter digestibility (DMD: 57.82%), total digestible nutrients (TDN: 54.76%) and comparatively gained 50 g more daily body weight.

India has approximately 600 million livestock, which requires almost 1000 million tons of hay or green fodder to sustain present level of productivity. However, as per estimates nearly 230 million tons of green fodder is available and the livestock has to struggle with the devastating scarcity of approximately 800 million tons of green fodder. The deteriorating animal health and their sustainability could also pose a potential threat for human existence and their livelihood. This necessitates the use of certain alternative options such as the agricultural crop residues and grasses (lignocellulosic biomass) as feed sources. If these are utilized judiciously this may provide enough energy and nutrients to the animals. However, high lignin content and lower digestibility and protein content and poor palatability of crop residues and grasses discourage their use as the sole animal feed. Lignin, being a cementing material in plant cell wall restricts the fullest accessibility of carbohydrates, the energy reserve, to the microorganisms inside the gut of ruminating animals. Despite employing various physical, chemical and physicochemical methods to remove lignin from the plant residues, currently the research efforts are focused on biological alternatives, especially because they are environmentally benign and widely acceptable.

Among various microorganisms known for lignin degradation, white-rot fungi (majorly basidiomycetes) have been adjudged most promising lignin degraders and have been largely studied for bioconversion of plant residues into nutritionally digestible animal feed under solid-state fermentation (SSF) conditions. However, these white-rot fungi have differential ability and mode to degrade lignin, either simultaneously or selectively. The white-rot fungi, which selectively degrade lignin without affecting much of the carbohydrates, and in turn expose protected and available carbohydrates to ruminants are prerequisite biological agent and especially considered well suited for animal feed development. Out of various selective lignin degrading fungi, only few have been studied in detail (Lentinus edodes, Pycnoporus cinnabarinus, Cereporiopsis subvermispora and Phlebia brevispora) at laboratory scale. To the best of our knowledge except Zadrazil and coworkers, who have demonstrated the ability of Pleurotus sp., to ferment straw in a 1.5 t capacity solid state bioreactor, large scale production of animal feed is scarcely been reported. However, they emphasized to conduct large scale animal feeding trials to further establish the suitability of the process of feed development.

Moreover, the majority of the white-rot fungi have been observed to degrade lignin considerably but at an expense of long fermentation period and thereby caused an undesirable reduction in carbohydrates and eventually digestibility of the fermented material. This does not only hamper the commercialization but also affects the economic viability of the process. Therefore, it is imperative to employ a fast growing and selective lignin degrading fungus for cost effective, nutrient rich and digestible animal feed development. For any animal feed developed, it is imperative to test it in cattle and before designing large scale in vivo animal trials of fermented feed it becomes priority to assess the quality and nutritional value of the feed through in vitro testing. Among different in vitro feed analyses, the gas production test hold specific importance, in which the amount of gas (CO₂ and CH₄) released, is measured when feeds are incubated in vitro with rumen liquor. The amount of gas produced...
is closely related to the digestibility and the energetic feed value of diets for ruminants\(^1\)\(^2\)\(^3\). The feeds with varied digestibility and nutrients are known to produce different volumes of gases within a stipulated time. This fact is taken in to consideration while designing diets for in vivo animal feeding trials to ensure the nutritional balance and adequacy of diet for animals. Moreover, in certain extended efforts, attempts have been made to replace either feed concentrate or grains from traditional animal diets with fermented wheat straw, especially to bring down the cost of animal feed\(^1\)\(^2\)\(^3\). However, to the best of our knowledge none of the study could be transformed into a wholesome process or technology. Therefore, there is a need to develop a wholesome/fermentation based process or technology to transform crop residue(s) in to nutritionally rich and digestible cattle feed.

This paper deals with developing a process for bioconversion of wheat straw into a digestible and nutrient rich animal feed with a selective lignin degrading fungus, *Crinipellis* sp. RCK-1, grown under solid-state fermentation for 5 days. The fermented feed produced, called here “Biotech Feed” has been evaluated in vitro followed by in vivo testing in buffalo calves, which has found to replace 50% grains from feed concentrate mixture.

**Results**

**Cell wall compositional changes of wheat straw.** *Crinipellis* sp. RCK-1 grew luxuriantly on wheat straw under solid state fermentation (SSF) conditions, which could be because of homogenous distribution of fungal pellets as an inoculum. The unfermented (control) wheat straw found to have (% w/w) ADF: 53.74, NDF: 83.10, hemicellulose: 29.36; lignin: 10.53; cellulose: 39.50 ash: 3.71 and 2.95% of crude protein. At 100 g level SSF, *Crinipellis* sp. RCK-1 caused higher degradation in lignin (28.26%) till 15th day of fermentation and consumed lesser cellulose (15%) along with a 48% decrease in hemicellulose content (Table 1). The plant cell wall degradation profile of *Crinipellis* sp. RCK-1 clearly showed that the fungus degraded lignin at a faster rate than cellulose on 10th day. The percent SSF efficiency, which is a measure of amount of lignin degradation at the cost of carbohydrate content loss, was found maximally increased on 10th day (Table 1). The fungal fermentation also caused a significant increase of 14.31% in total crude protein content in the Biotech Feed till 10th day of incubation. More interestingly, upon scale up of Biotech Feed production process in Koji room (500 g substrate in each tray), *Crinipellis* sp. RCK-1 exhibited similar growth and substrate colonization and the substrate degradation was comparable to that was observed at 100 g substrate level SSF in smaller trays. However, a slight increase in carbohydrate (cellulose and hemicellulose) degradation was observed in scale up experiments, but lignin degradation remained almost same (16.06%) till 9th day with a maximal increase in crude protein by 40.81% (Table 1). Moreover, *Crinipellis* sp. RCK-1 degraded a fair amount of lignin (~7%) till 5 day irrespective of the scale of experiment while, preventive much of cellulose degradation and concurrently increased the crude protein (up to 15–18%).

**In vitro evaluation of fermented feed.** In vitro gas production test was conducted to evaluate the changes in digestibility and nutrients of Biotech feed as an effect of fungal fermentation. The oven dried Biotech feed fermented for 5, 10 and 15 days along with the unfermented wheat straw (WS) were tested for in vitro gas production. True degradable organic matter (TDOM), microbial biomass production (MBP), organic matter digestibility (OMD) and metabolizable energy (ME) of 5 day Biotech Feed (BT5) was significantly higher (P<0.05) as compared to 10th (BT10) and 15th day (BT15) Biotech Feed. BT5 exhibited a significant improvement in OMD and ME of ~30% when compared with unfermented straw (Table 2). However, no significant difference was found in ME, OMD, MBP and TDOM of BT10 and BT15, which further suggested carrying out SSF of wheat straw till 5 day only. OMD and ME, however was found to be improved up to 10–12% in comparison to the unfermented wheat straw on 15 day of incubation, but 5 day fermented Biotech Feed (BT5) was found to be most suitable for in vivo animal testing due to its higher energy content and available carbohydrate based nutrients.

The animal diets were also characterized to assess their nutritional quality in terms of their adequacy to meet out the animal maintenance and potential to replace certain amount of grains from traditional feed concentrates. Keeping in view the nutrient balance, four diets i.e. T1 control (WS + concentrate mixture), T2 (5 day Biotech Feed + concentrate mixture), T3 (5 day Biotech feed + 50% grain replaced from concentrate mixture), T4 (5 day Biotech Feed + 100% grain replaced from concentrate mixture) were tested. The organic matter digestibility was significantly higher in T2 and T3 diets as compared to T1 and T4. A same trend was found for gas production (GP), TDOM and MBP. While, T2 had maximum ME (MJ/kg DM) followed by T3, T4 and T1 and the difference among the four treatments was found significantly differing (Table 3). The in vitro analysis suggested that 50% grain can be replaced from concentrate mixture in Biotech Feed based diet (T3), without compromising with the energy and nutrients availability of the diet. However, replacement of 100% grains from concentrate (T4 diet) reduced OMD 40.65% and ME by 5.47 MJ/kg DM and hence was not included in in vivo digestibility trial further. The partitioning factor (PF) was found reduced in diets as compared to control which in turn exhibits improved available energy in fermented diets (Table 3).

**Amino acid analysis of fermented wheat straw.** Amino acid analysis of unfermented and *Crinipellis* sp RCK-1 5 day fermented feed (BT5) revealed that fungal fermentation has considerably improved the individual amino acid content. Among various essential amino acids, Arg, Thr, Ileu and Leu were found increased essential amino acids, Arg, Thr, Ileu and Leu were found increased

| Table 1 | Cell wall composition (% w/w) of control and fungal treated wheat straw (*Crinipellis* sp. RCK-1) under solid state fermentation (SSF) condition |
|---|---|---|---|---|---|---|
| Fermentation level (Substrate weight) | Day | Hemi-cellulose | Lignin | Cellulose | % Eff. of SSF | Crude protein |
| Unfermented control | 0 | 29.36 ± 1.72 | 10.53 ± 0.013 | 39.50 ± 0.44 | - | 2.95 ± 0.116 |
| 100 g | 5 | 23.56 ± 1.55 (−19.76) | 9.84 ± 0.002 (−6.56) | 37.33 ± 1.15 (−5.48) | 8.67 | 3.418 ± 0.147 (±15.87) |
| | 10 | 20.87 ± 2.27 (−28.93) | 8.60 ± 0.002 (−18.36) | 35.43 ± 0.47 (−10.31) | 15.38 | 3.372 ± 0.129 (±14.31) |
| | 15 | 15.24 ± 0.62 (−48.08) | 7.55 ± 0.014 (−28.26) | 33.21 ± 0.59 (−15.92) | 14.59 | 3.369 ± 0.201 (±14.19) |
| | 5 | 23.74 ± 1.37 (−19.16) | 9.78 ± 0.007 (−7.16) | 36.71 ± 0.84 (−7.06) | 8.98 | 3.494 ± 0.198 (±18.43) |
| | 10 | 21.49 ± 2.17 (−26.81) | 9.34 ± 0.006 (−11.26) | 33.94 ± 1.03 (−14.07) | 8.82 | 3.50 ± 0.126 (±18.64) |
| | 15 | 19.61 ± 1.36 (−33.2) | 8.84 ± 0.001 (−16.06) | 33.09 ± 1.57 (−16.22) | 10.46 | 4.154 ± 0.115 (±40.81) |

\(*\) Sign denotes the standard deviation (S.D.) and values in parenthesis represent % degradation of different components with respect to unfermented control (+ % increase and - % decrease).

% efficiency of SSF is a derived ratio from the values of these components ESSF = (loss of lignin/loss of hemicellulose + cellulose) * 100.
up to 33.33, 34.30, 31.58 and 30.77%, respectively (Table 4). Since there was an increase in total crude protein content, an increase in non essential amino acids was also noticed. Cys was not present in wheat straw, whereas BT5 was found to be maximally enriched with ergosterol content (292.17 g/gds). Unfermented wheat straw contained 19.44 μg/g dry substrate of ergosterol content, while BT5 was found to be maximally enriched with ergosterol content (292.17 μg/gds).

### Analysis of fungal biomass enrichment through ergosterol estimation.

Crinipellis sp RCK-1 5 day fermented feed (BT5) was found to have significantly higher amount of fungal cell mass. Fungal biomass was quantified in term of total ergosterol content in unfermented and BT5 straw. Unfermented wheat straw contained 19.44 μg/g dry substrate (gds) of ergosterol content, while BT5 was found to be maximally enriched with ergosterol content (292.17 μg/gds).

#### Structural characterization of fungal fermented feed by electron microscopy.

Structural transformation in wheat straw due to fermentation was studied by SEM and TEM. Scanning electron micrographs of unfermented and fermented wheat straw revealed that there was a heavy fungal attack on wheat straw surface and extended to vascular bundles and adjoining cells i.e. parenchymatic cells (Fig. 1a, b). The structural matrix of wheat straw was observed to be completely distorted and fungal hyphae were clearly visible within the fermented wheat straw (Fig. 1b). The hyphae of Crinipellis sp. RCK-1 appeared to be penetrated deep within the wheat straw and dismantled the original structure (Fig. 1c). Whereas, SEM pictures of unfermented wheat straw showed clear visible organized structure of cell lumina and intact vascular bundles (Fig. 1a).

TEM figures of unfermented wheat straw possessed intact cell wall structure as electron dense region and an organized cork like structure of the digestible fiber, protein, lipid, and carbohydrate components of the digestible fiber, protein, lipid, and carbohydrate components of the straw cell walls have shown either complete degradation of lignin-rich middle lamellae, between S1 and S2 layers or selective degradation of lignin from cell walls of wheat straw (Fig. 1e, f). A clear reduction in electron dense region was observed within the cell walls and aggregates of lignin residuals were seen in cellular skeleton. Moreover, fungal hyphae were also observed in cell lumina (Fig. 1f).

### In vivo digestion trial in buffalo calves.

Chemical composition of feed and fodder. All the three diets were balanced for their nutritional value i.e. crude protein and organic matter. The ingredients of diets i.e. wheat straw, Biotech Feed, concentrate mixture and green fodder were analyzed for their proximate composition. The crude protein of the Biotech Feed increased from 3.15 to 3.65%, while neutral detergent fiber (NDF) and Acid detergent fiber (ADF) content in Biotech Feed were decreased from 83.42, 56.86 to 78.40, and 53.60%, respectively, as compared to unfermented wheat straw. The ash content of Biotech Feed was also found increased from 7.40 to 10.20% compared to control wheat straw.

Voluntary intake of nutrients. Dietary group T2 recorded highest dry matter intake in terms of kg body weight (BW)% and g kg⁻¹ metabolic body weight (W⁻⁰.₇⁵) followed by T1 and T3. The values of digestible dry matter and digestible organic matter intake (kg BW% and g kg⁻¹ W⁻⁰.₇⁵) were significantly higher in Biotech Feed based diets T2 and T3 in terms of kg BW% and g kg⁻¹ W⁻⁰.₇⁵ (Table 5).

Digestibility coefficients of nutrients. Total tract digestibility was significantly (P<0.05) improved among calves fed on Biotech feed based diets (T2 and T3) (Table 6). The DM and OM digestibility were 52.07 and 54.52%, respectively, in the wheat straw based diet T1, whereas they were found as 57.82 and 60.54% in T2; 57.47 and 59.83% in T3 diets, respectively. The digestibility of NDF, ADF, ether extract (EE) and roughage were significantly affected by the treatments except crude protein digestibility, which was found not significantly differing. The straw dry matter digestibility (DMD) was significantly higher in Biotech Feed based diets (T2 and T3) than with control group. The average values of roughage dry matter digestibility were 32.22, 42.61 and 39.16% in T1, T2 and T3, respectively.

Nutrient density. The total digestible nutrient is known as the sum of the digestible fiber, protein, lipid, and carbohydrate components

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| Sample                        | TDOM (mg) | MBP (mg) | OM digestibility (%) | ME (MJ/kg DM) |
|-------------------------------|-----------|----------|----------------------|---------------|
| Wheat straw                   | 136.82ab  | 107.11bc | 28.05bc              | 4.06a         |
| 5 days old Biotech Feed (BT5) | 162.15a   | 110.95a  | 36.74a               | 5.38b         |
| 10 days old Biotech Feed (BT10)| 135.84a  | 98.07b   | 31.31a               | 4.56a         |
| 15 days old Biotech Feed (BT15)| 138.70a  | 101.80a  | 30.98a               | 4.50a         |
| SEM ±                         | 3.34      | 2.68     | 1.27                 | 0.378         |

Values bearing different superscripts (a-b) in a column differ significantly (P < 0.05), while same superscripts (a-d, ab) denotes non significant difference. Overall SEM ± has been given by first calculating sample mean then calculating overall mean in a column (j) for an individual parameter in different groups.

**Table 2** | In vitro gas production profile of fermented Biotech Feed

| Sample                        | IVGP/200mg | TDOM/200mg | TDOM% | OMD% | MBP/200mg | EMP | PF | ME (MJ/kg DM) |
|-------------------------------|------------|------------|-------|------|-----------|-----|----|---------------|
| T1 Control (WS + Conc. mixture)| 25.27      | 106.72    | 53.36 | 40.21 | 57.37     | 53.76 | 4.25 | 5.45          |
| T2 (Biotech Feed + Conc. mixture)| 30.05      | 120.22   | 60.11 | 44.09 | 65.69     | 54.64 | 4.01 | 6.00          |
| T3 (Biotech Feed + 50% Grain replaced from conc. mixture)| 29.53      | 118.41   | 59.20 | 43.54 | 65.06     | 54.94 | 4.01 | 5.91          |
| T4 (Biotech Feed + 100% grain replaced from conc. mixture)| 26.10      | 108.28   | 54.14 | 40.65 | 59.83     | 55.24 | 4.15 | 5.47          |
| SEM ±                         | 0.54       | 1.55      | 0.81 | 0.45 | 0.91      | 0.20 | 0.04 | 0.07          |

Values bearing different superscripts (a-d) in a column differ significantly (P < 0.05), while same superscripts (a-d, ab) denotes non significant difference. Overall SEM ± has been given by first calculating sample mean then calculating overall mean in a column (j) for an individual parameter in different groups.
of a feedstuff or diet, which is directly related to digestible energy. The total digestible nutrient (TDN) % in Biotech Feed diet based groups (T2 and T3) was found significantly \((P<0.05)\) higher (Table 7). Whereas, the digestible crude protein (DCP) is largely determined by subtracting unavailable protein from crude protein. The digestible crude protein content was found to be improved in T2 and T3 diets non significantly higher when compared with the control diet (T1).

### Blood biochemical analysis

Blood biochemical analysis showed that there were non-significant differences in packed cell volume (PCV), haemoglobin (Hb), total protein, albumin and serum creatinine, but significant difference were found in blood urea nitrogen (BUN) and cholesterol level (Table 7). Low levels of cholesterol and alkaline phosphatase in T3 group is an indicative of better excretory function by calves fed Biotech Feed based diet. Higher level of BUN in the T2 and T3 groups indicated the higher rate, is known to play only a minor role in supplying nutrients in the cattle rumen\(^4\). Earlier, various fungi i.e. Pleurotus sp., T. versicolor, C. subvermispora, Ganoderma sp., B. adusta, L.edodes and P. brevispora have been studied for bioconversion of wheat straw to digestible animal feed. But the majority of them took quite a long period in degrading either similar or higher amounts of lignin compared to Crinipellis sp. RCK-1, i.e. Pleurotus sp., (~37% in 30 days) \(T.\) versicolor (~31% in 30 days), C. subvermispora (62% in 49 days), Ganoderma sp. (~11% in 49 days), B. adusta (42% in 49 days), L.edodes (58.9% in 49 days) and P. brevispora (45.4% in 49 days)\(^5,6,7\).

### Discussion

Crinipellis sp. RCK-1 was found to degrade lignin in wheat straw faster on 5\(^{th}\) day than cellulose and hemicellulose. However, the hemicellulose content, which was observed to degrade at a slightly higher rate, is known to play only a minor role in supplying nutrients in the cattle rumen\(^8\). Earlier, various fungi i.e. Pleurotus sp., T. versicolor, C. subvermispora, Ganoderma sp., B. adusta, L.edodes and P. brevispora have been studied for bioconversion of wheat straw to digestible animal feed. But the majority of them took quite a long period in degrading either similar or higher amounts of lignin compared to Crinipellis sp. RCK-1, i.e. Pleurotus sp., (~37% in 30 days) \(T.\) versicolor (~31% in 30 days), C. subvermispora (62% in 49 days), Ganoderma sp. (~11% in 49 days), B. adusta (42% in 49 days), L.edodes (58.9% in 49 days) and P. brevispora (45.4% in 49 days)\(^5,6,7\).

### Table 4 | Amino acid content of unfermented and fungal fermented wheat straw

|                      | Control | CR 5 | % Increase |
|----------------------|---------|------|------------|
| **Non-essential amino acids** |         |      |            |
| Asp                  | 0.207   | 0.260 | 20.51      |
| Glu                  | 0.480   | 0.653 | 26.53      |
| Ser                  | 0.120   | 0.187 | 35.71      |
| Gly                  | 0.127   | 0.287 | 55.81      |
| Ala-pro              | 0.167   | 0.233 | 28.57      |
| Tyr                  | 0.100   | 0.073 |            |
| Cys                  | 0.000   | 0.040 | 100.00     |
| **Essential amino acids** |         |      |            |
| Val                  | 0.133   | 0.127 |            |
| Met                  | 0.267   | 0.080 |            |
| Ileu                 | 0.087   | 0.127 | 31.58      |
| Leu                  | 0.060   | 0.087 | 30.77      |
| Phe                  | 0.000   | 0.000 | 0.00       |
| Lys                  | 5.800   | 5.613 |            |
| His                  | 0.013   | 0.013 | 0.00       |
| Arg                  | 0.067   | 0.100 | 33.33      |
| Thr                  | 0.153   | 0.233 | 34.29      |
| Total AA             | 7.780   | 8.113 | 4.28       |

| g/100 gds | Control | CR 5 | % Increase |
|-----------|---------|------|------------|
| Non-essential amino acids |         |      |            |
| Asp | 0.207   | 0.260 | 20.51      |
| Glu | 0.480   | 0.653 | 26.53      |
| Ser | 0.120   | 0.187 | 35.71      |
| Gly | 0.127   | 0.287 | 55.81      |
| Ala-pro | 0.167   | 0.233 | 28.57      |
| Tyr | 0.100   | 0.073 |            |
| Cys | 0.000   | 0.040 | 100.00     |
| Essential amino acids |         |      |            |
| Val | 0.133   | 0.127 |            |
| Met | 0.267   | 0.080 |            |
| Ileu | 0.087   | 0.127 | 31.58      |
| Leu | 0.060   | 0.087 | 30.77      |
| Phe | 0.000   | 0.000 | 0.00       |
| Lys | 5.800   | 5.613 |            |
| His | 0.013   | 0.013 | 0.00       |
| Arg | 0.067   | 0.100 | 33.33      |
| Thr | 0.153   | 0.233 | 34.29      |
| Total AA | 7.780   | 8.113 | 4.28       |

gds- gram dry substrate.
Control – Unfermented wheat straw.
CRS- Crinipellis sp. RCK-1, 5 day fermented wheat straw.

Figure 1 | Scanning electron micrograph of unfermented (a- 100 \(\mu m\)) and Crinipellis sp RCK-1 5 day fermented wheat straws (b) at 100 \(\mu m\) (c) 10 \(\mu m\) and transmission Electron Micrograph showing selective lignin degradation, (d–e) unfermented and (f) Crinipellis sp RCK-1, 5 day fermented wheat straw. Abbreviations: CC- Cell corner, S1–S2- layers, ML- Middle lamellae.

### Table 5 | Voluntary intake of different nutrients in calves under different treatments

| Particulars | T-1 | T-2 | T-3 | SEM ± |
|-------------|-----|-----|-----|-------|
| DMI (kg day\(^{-1}\)) | 3.27\(^a\) | 3.74\(^a\) | 2.97\(^a\) | 0.184 |
| DMI (kg BW\%) | 2.80\(^a\) | 2.95\(^a\) | 2.92\(^a\) | 0.046 |
| DMI (g kg\(^{-1}\)W\(^{0.75}\)) | 91.95\(^a\) | 98.21\(^b\) | 93.03\(^a\) | 1.188 |
| DMI (kg day\(^{-1}\)) | 2.99\(^a\) | 3.38\(^a\) | 2.65\(^a\) | 0.169 |
| DMI (kg BW\%) | 2.57\(^a\) | 2.66\(^a\) | 2.61\(^a\) | 0.040 |
| DMI (g kg\(^{-1}\)W\(^{0.75}\)) | 84.29\(^a\) | 88.82\(^a\) | 82.99\(^a\) | 1.090 |
| DDMI (kg day\(^{-1}\)) | 1.71\(^a\) | 2.16\(^a\) | 1.71\(^a\) | 0.110 |
| DDMI (kg BW\%) | 1.46\(^a\) | 1.70\(^a\) | 1.68\(^ab\) | 0.042 |
| DDMI (g kg\(^{-1}\)W\(^{0.75}\)) | 47.91\(^a\) | 56.79\(^a\) | 53.47\(^ab\) | 1.275 |
| DOMI (kg day\(^{-1}\)) | 1.64\(^a\) | 2.04\(^a\) | 1.59\(^a\) | 0.105 |
| DOMI (kg BW\%) | 1.40\(^a\) | 1.61\(^a\) | 1.56\(^ab\) | 0.039 |
| DOMI (g kg\(^{-1}\)W\(^{0.75}\)) | 40.07\(^a\) | 45.03\(^ab\) | 48.87\(^bc\) | 2.700 |

Values bearing different superscripts (a–c, bc) in a row differ significantly \((P<0.05)\), while same or common superscripts (a–c, ab) denotes non significant difference. Overall SEM ± has been given by first calculating sample mean then calculating overall mean in a row \((\pm)\) for an individual parameter in different groups. DMI, dry matter intake; OMI, organic matter intake; DDMI, digestible dry matter intake; DOMI, digestible organic matter intake, BW, body weight; \(W^{0.75}\), metabolic body weight.
Prolonged incubation periods have already been reported to be found associated with simultaneous degradation of cellulose and hemicellulose eventually causing an undesirable decrease in dry matter digestibility of the fermented substrate. Since the selective ligninolysis is characterized by a higher positive correlation between total organic matter loss and lignin loss as compared to polysaccharides degradation\textsuperscript{43}. Hence, the present study advocated carrying out the organic matter loss and lignin loss as compared to polysaccharides during fungal fermentation\textsuperscript{30}. Contrary to that, consistent increase in digestibility with increase in fungal degradation of lignocellulosic components during prolonged incubations (30–60 days) has also been reported by several workers\textsuperscript{6,35,36}. In vitro analysis of Biotech feed based diets depicted that even after reduction of 50% grains, which in turn reduced 0.34 Mcal/kg dry matter from concentrate mixture, no significant decrease was observed in OM digestibility, metabolizable energy and microbial biomass production. This clearly indicated that Biotech feed has the potential to replace 50% grains from traditional feed concentrates and paved the way for in vivo evaluation of feed.

However, based on various studies it is widely accepted that the extent of lignin degradation and increase in in vitro digestibility largely depend on the fungus and incubation conditions and hence can not be generalized\textsuperscript{47,48}. Recently, Tuyen et al\textsuperscript{48} have also shown that removing lignin alone does not always improve the in vitro gas production in the glass syringe test or in vitro digestibility and a weak correlation (\( r=0.47 \)) has been observed between them. Moreover, some studies have also reported a strong and significant correlation between the increase in gas production and lignin to cellulose ratio (percent SSF efficiency in present study) and inclusion of hemicellulose loss in the analysis has shown to increase the goodness of fit of the equation\textsuperscript{44}. Furthermore, reduced partitioning factor (PF) in diets as compared to control is also a good sign of improved available energy in fermented diets.

Amino acids are required for protein synthesis for maintenance, growth and productivity of animal\textsuperscript{41}. Essential and non-essential amino acids are generally discriminated on the basis of metabolic capability of animal to synthesize it or not. Rumen microbes are capable of synthesizing amino acids for microbial protein synthesis only when sufficient carbon source (majorly from dietary carbohydrate), non- protein nitrogen (from inorganic supplementation) and inorganic sulphur are available\textsuperscript{7}. It was thereby inferred, that fermentation of wheat straw by \textit{Crinipellis sp.} RCK-1 provided sufficient nutrients to support microbial protein synthesis, eventually causing an increase in essential amino acids as well. The better assim-

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|c|}
\hline
\textbf{Treatment (diets)} & \textbf{DM} & \textbf{Roughage} & \textbf{OM} & \textbf{CP} & \textbf{EE} & \textbf{NDF} & \textbf{ADF} \\
\hline
\textbf{T\textsubscript{1}} & 52.07\textsuperscript{a} & 32.22\textsuperscript{a} & 54.52\textsuperscript{a} & 68.96\textsuperscript{a} & 72.36\textsuperscript{a} & 45.81\textsuperscript{a} & 36.69\textsuperscript{a} \\
\textbf{T\textsubscript{2}} & 57.82\textsuperscript{a} & 42.61\textsuperscript{a} & 60.54\textsuperscript{a} & 74.72\textsuperscript{a} & 75.15\textsuperscript{a} & 53.33\textsuperscript{ab} & 51.42\textsuperscript{a} \\
\textbf{T\textsubscript{3}} & 57.47\textsuperscript{ab} & 39.16\textsuperscript{b} & 59.83\textsuperscript{ab} & 69.97\textsuperscript{b} & 67.25\textsuperscript{b} & 53.47\textsuperscript{b} & 47.42\textsuperscript{b} \\
\textbf{SEM\pm} & 0.848 & 1.492 & 0.862 & 2.154 & 1.115 & 1.1198 & 2.031 \\
\hline
\end{tabular}
\caption{Digestibility of different nutrients (%) in calves under different treatments}
\end{table}

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|}
\hline
\textbf{Nutritive Value (%)} & \textbf{T\textsubscript{1}} & \textbf{T\textsubscript{2}} & \textbf{T\textsubscript{3}} & \textbf{SEM \pm} \\
\hline
\textbf{No. of Animals} & 4 & 4 & 4 & - \\
\textbf{Experimental period (days)} & 90 + 7 = 97 & 90 + 7 = 97 & 90 + 7 = 97 & - \\
\textbf{Total digestible nutrient} & 49.97\textsuperscript{a} & 54.76\textsuperscript{a} & 53.38\textsuperscript{a} & 0.665 \\
\textbf{Digestible crude protein} & 8.42\textsuperscript{b} & 8.77\textsuperscript{a} & 8.80\textsuperscript{b} & 0.254 \\
\textbf{Initial live body weight (kg)} & 84.50\textsuperscript{c} & 85.00\textsuperscript{b} & 82.25\textsuperscript{a} & 4.029 \\
\textbf{Final live body weight (kg)} & 116.53\textsuperscript{a} & 121.53\textsuperscript{a} & 112.98\textsuperscript{a} & 4.503 \\
\textbf{Total gain (kg)} & 32.02\textsuperscript{a} & 36.52\textsuperscript{a} & 30.72\textsuperscript{a} & 1.111 \\
\textbf{Av. Daily Gain (g/d)} & 355.83\textsuperscript{a} & 405.83\textsuperscript{a} & 341.39\textsuperscript{a} & 12.364 \\
\textbf{Feed Conversion (kg DMI/kg gain)} & 9.16\textsuperscript{a} & 9.18\textsuperscript{a} & 8.76\textsuperscript{a} & 0.340 \\
\textbf{PCV (Packed cell volume)} & 33.00\textsuperscript{a} & 33.25\textsuperscript{a} & 34.75\textsuperscript{a} & 0.742 \\
\textbf{Haemoglobin (g/dl)} & 9.75\textsuperscript{a} & 10.25\textsuperscript{a} & 10.13\textsuperscript{a} & 0.875 \\
\textbf{Total protein (g/dl)} & 7.45\textsuperscript{a} & 7.95\textsuperscript{a} & 7.83\textsuperscript{a} & 0.160 \\
\textbf{Albumin (g/dl)} & 7.20\textsuperscript{b} & 8.00\textsuperscript{b} & 7.73\textsuperscript{b} & 0.064 \\
\textbf{BUN (blood urea nitrogen) (mg/dl)} & 7.62\textsuperscript{a} & 16.83\textsuperscript{b} & 16.25\textsuperscript{b} & 1.483 \\
\textbf{Serum creatinine (mg/dl)} & 1.10\textsuperscript{b} & 1.13\textsuperscript{a} & 1.15\textsuperscript{a} & 0.059 \\
\textbf{Alkaline phosphatase (U/L)} & 325.00\textsuperscript{a} & 293.75\textsuperscript{a} & 241.50\textsuperscript{a} & 34.675 \\
\textbf{Cholesterol (mg/dl)} & 153.43\textsuperscript{a} & 133.23\textsuperscript{ab} & 93.43\textsuperscript{c} & 8.991 \\
\hline
\end{tabular}
\caption{Nutrient density (%), growth performance and blood biochemical profile of calves fed under different treatments}
\end{table}
ulation of nutrients and adequate supply of amino acids supported not only the maintenance but also resulted in an improved body weight gain among animals fed on fermented diets. Supplementation of the essential amino acid is already been emphasized to achieve a low cost feed formulation\(^\text{43}\).

Analysis of ergosterol as fungal biomass index holds specific importance majorly because of the rare 5–7 double bonding based sensitive assay in UV range in neutral lipid extracts since this bonding is rarely reported in major sterols of plants\(^32\). Fungal biomass estimation is a major problem while determining fungal growth especially in solid state fermented products, since the biomass grows and remains trapped inside the substrate. Ergosterol content has been established a reliable indicator of fungal growth under SSF conditions and it majorly represents the live biomass\(^32\). The fungal biomass could have been determined in terms of chitin, however, this method has faced a number of criticism specially, the tendency of the chitin content in mycelium to vary with age and the substrate specific interference with the assay\(^32,33\). While, ergosterol, the predominant sterol of most fungi, found almost exclusively in membranes of living fungal cells, is not produced in significant quantities by green plants and hence can be used as an index of fungal colonization\(^32,33\). Moreover, the ergosterol method is more sensitive and easier and takes only 5–6 h and has the tenacity to be scaled up to cope with large numbers of samples within the same time scale\(^34\). Therefore, fungal biomass was estimated employing ergosterol based method.

Enrichment of fermented feed with enormous fungal biomass displays development of an efficient bioprocess of wheat straw bioconversion, providing an environment for a luxuriant growth of the fungus within substrate eventually producing more digestible and nutrient rich cattle feed.

Within 5 days of incubation, *Crinipellis* sp. RCK-1 efficiently colonized and degraded straw as has been clearly demonstrated through surface micrographs of the unfermented and fermented wheat straw. The SEM micrographs were also taken at the edges of wheat straw to provide a better view of degradation and to capture the sites of distortion. Middle lamella and S2 layer showed a clear separation of tissues as has been observed by Berrocal et al\(^42\) during biological upgrading of wheat straw under SSF conditions by *Streptomyces cyanus*. The degradation of lignin rich middle lamella and cell corners have supported visually that the lignin was degraded in the wheat straw\(^41\). Presence of fungal hyphae inside the cell lumina further confirmed the deep penetration by the fungus and its progressive degradation of the wheat straw. However, unlike various other studies, *Crinipellis* sp. RCK-1 has shown the potential to selectively degrade lignin irrespective of the tissue ranging from most lignified xylem and sclerenchyma to less lignified parenchyma\(^41-42\). This observation clearly demonstrated the robust nature of *Crinipellis* sp. RCK-1 and its potential to convert poor quality wheat straw into nutritive animal feed without affecting much of the cellulose and hemicellulose portions of the cell walls. The electron dense residuals of lignin termed here as lignin aggregates have also been demonstrated earlier by many workers and they are considered to be a common characteristic of an advanced stage of white rot decay\(^41-42\).

*In vitro* gas production (IVGP) has been reported to be advantageous over *in vivo* methods for estimation of nutrient availability in cattle feed, being less expensive, less time consuming, requiring small amounts of samples allowing a better quantification of nutrient utilization and accuracy in describing digestibility. However, to ensure the suitability of fermented feed, *in vivo* digestion trials are essential and hence it was attempted to establish the nutritive potential of fermented straw as cattle feed through *in vivo* feeding\(^28,40\). Such a long period of 90 days was chosen to ensure the consistency in animal performance, minimizing the fluctuation in feed intake and to determine any refusal by the calves. Only after the 90 days when, feed intake, palatability and left over (if any) became constant, 7 days digestion trial was carried out.

The results were instrumental with various *in vivo* animal feeding trials carried out under similar conditions. In accordance to the present study, an increase in DMI (% of body weight/day) has been observed in lambs by Calazada et al\(^28\), which were fed with *P. sajor caju* fermented wheat straw. In a similar feeding trial using Murrah male buffalo calves, an attempt was made to replace wheat straw/rice straw (in control diet) with *Pleurotus* treated spent wheat straw at a rate of 0% (T1), 50% (T2) and 100% (T3)\(^36\). It was observed that DMI (kg/100 kg body weight) was found to be increased as 1.63 kg/100 kg body weight in T2 diet fed calves when compared with T1 (1.56 kg/100 kg body weight) in case of wheat straw, while it remained unchanged in animal fed with rice straw based diets. Similarly, during an *in vivo* feeding of Simmental heifers, an attempt was made to replace 26% (group II) and 44% (group III) maize grains with *P. ostreatus* mushroom compost wheat straw, but a sudden decrease in average daily gain was observed from 1150 g/d in control group (I) to 1140 and 990 g/d, respectively\(^37\). Likewise, regular fluctuations have also been noticed by Fazaeli et al\(^29\) in parameters such as DMI (g/d), OMI (g/d), DMI (g/kg BW\(^{0.75}\)), OMI (g/kg BW\(^{0.75}\)), digestible dry matter intake (DDMI) and digestible organic matter intake (DOMI) g/d and DDMI and DOMI (g/kg BW\(^{0.75}\)), when male sheep were fed with fermented wheat straw treated with different *Pleurotus* strains.

In this study, the highest values for nutrients digestibility were observed in T2, which was significantly higher than other treatments. Similar observation for nutrient digestibility has been noticed by many other workers while feeding fungal fermented diets to the monogastric animals and ruminants i.e. lambs, goats, buffalo calves, Simmental heifers, sheep and Hanwoo steers\(^27,28,31,32,34,35\). However, non-significant improvement in *in vivo* digestibility for certain nutrients i.e. crude protein and ether extract has also been reported by Okano et al\(^39\) and Shirivastava et al\(^40\). However, in contrast to our findings a decrease in TDN was observed when chemically (urea) treated (UTRS) and fungal treated (FTRS) rice straw were fed to cross bred goats i.e. 51.28 and 38.38%, respectively\(^41\). This decrease was linked with a higher dry matter loss and loss of potential energy materials by *Coprinus fimetarius* (currently known as *copepinopsis cinerea*) during prolonged fungal fermentations. The increase in TDN content in our study is a clear indicator of improved and intact availability of energy substances in Biotech Feed.

The gradual increase in DCP in the T2 and T3 dietary groups represented an improved assimilation of nitrogen by ruminants as has been reported by Kakkar and Dhandha\(^39\), where replacement of 50 and 100% plain wheat straw with *Pleurotus* sp. fermented straw in diets (T2 and T3, respectively) improved DCP. Interestingly, in the present study the T3 group, where 50% grains from concentrate mixture have been replaced by Biotech Feed, exhibited highest DCP as 8.8% and improved TDN, which eventually advocated the possibility of using T3 diet as a sole ration to the ruminants in future. Our present results are consistent with our earlier report, where TDN and DCP were found increased upon feeding *Ganoderma* sp. rckk02 fermented wheat straw based diet to crossbred goats\(^46\). Growth rate in terms of average daily gain (ADG) of calves was also found effectively improved upon feeding Biotech Feed. In a similar study Dey et al\(^43\) have reported improvement in ADG, when *Orpinomycyss* sp., an anaerobic gut fungus belonging to the phylum Neocallimastigomycota, was administered at the rate of 106 CFU/ml/calf/week (diet T2) in addition to wheat straw + concentrate feed based diet (T1). The improved ADG among calves fed on diet T2 (709 g/d) compared to diet T1 (614 g/d) was attributed to the better utilization of fermented wheat straw because of the availability of more digestible carbohydrates\(^45,46\). Likewise, Salman et al\(^45\) have also shown that feeding of fermented sugar beet pulp supplemented diet at the rate of 0.6% (T3) and 0.9% (T4) to goats, significantly improved ADG up to 95.25 and 104.83 gram/hour/day (g/h/d), respectively, compared to a control diet (T1, unsupplemented) i.e. 88.58 (g/h/d). However, their study also clearly revealed that at the lower rate of supplementa-
tion i.e. 0.3% (T2), ADG could not be improved, as in case of our study.

In vivo feeding trials pertaining to grain replacement have not been much successful in past and a decrease in ADG has also been reported, which could largely be due to the problem of palatability, reduced feed conversion and animal refusal. Moreover, different to what has been much successful in past and a decrease in ADG has also been reported, which could largely be due to the problem of palatability, decreased feed conversion is undesirable and is known to have adverse effects on health of calves. The present study showed that no major and significant differences were noticed in blood metabolites among different dietary groups.

Inoculum development. Preparation of pellets in fungal shake flask (250 ml). The fungal pellets so developed were used as seed inoculum for further large scale inoculum production.

Preparation of fungal pellets in shake flask (2 L). Large scale inoculum production was carried out in 2 L shake flasks, each having 1 L of MEB sterilized at 121 °C (15 psi) inoculated with pellets developed as described above as seed inoculum and incubated at 30 °C for 72 hours and pellets thus developed were used as inoculum for large scale (500 g wheat straw in each tray) development of Biotech Feed.

Solid State Fermentation (SSF) of wheat straw. Solid State Fermentation (SSF) of wheat straw. The fungal pellets so developed were used as seed inoculum for further large scale inoculum production.

Scale up of solid state fermentation (SSF). Biotech Feed production was carried out by fermenting 50 kg of wheat straw in stainless steel trays (50.80 × 50.80 cm) inoculated with fungal pellet inoculum 1% (w/w on dry weight basis) and incubated for 5 days in a Koji room (Lignocellulose Biotechnology Lab, University of Delhi South Campus New Delhi, India). Koji room (3.04 × 3.04 Meter) is an especially designed fermentation facility equipped with controlled humidity (70 ± 5%) and temperature (30 ± 2 °C) and sterility through sterile air circulation system with HEPA filters (0.3 μm). Each single batch of solid state fermentation was comprised of 100 trays and each tray contained 500 g of wheat straw. Thus, each batch of SSF contained 50.0 kg of wheat straw and in all five such batches of fermentation were carried out in the lab scale experiment for each incubation period i.e. 5, 10 and 15 days. The trays were incubated at 30 °C for 5, 10 and 15 days. The trays without fungal pellets were used as control. The trays harvested after various fermentation periods were oven dried until it achieved constant weight and processed further for estimation of cell wall composition, proximate content and in vitro digestibility.

In vitro gas production test of Biotech Feed. The in vitro and in vivo trials were performed strictly as per NRC guidelines and with due permission of the Institutional Ethical committee. The in vitro digestibility of Biotech Feeds was analyzed at Indian Veterinary Research Institute (IVRI), Bareilly, UP, India. Two male buffaloes of 1-1.5 years age were used, which were surgically operated for fixing rumen fistula and tannin included standard diet for collecting fresh rumen liquor for studying in vitro gas production. The degradability of dry matter, organic matter and neutral detergent fibre was estimated following the method of Menke and Steinhauss. The samples were introduced into a calibrated glass syringe with buffered rumen liquor medium and inoculated at 39 °C. The gas produced was recorded from the glass syringe at the end of 24 hour incubation. Other parameters like organic matter digestibility (OMD), metabolizable energy (ME), short chain fatty acids (SCFAs), partitioning factor (PF) and microbial biomass (MBP) were calculated as described earlier. The PF is the amount (mg) of truly digested organic matter (TDOM) divided by the amount (mg) of metabolizable energy (ME) of the feed. It is generally taken as an index of the OMD between microbial biomass and fermentation gases produced during fermentation. Thus, the higher PF means that a higher proportion of TDOM is used for synthesis of microbial biomass. This has implications in rumen feeding.

Methods

Microorganism. Crinipellis sp. RCK-1 obtained from fungal culture collection of Lignocellulose Biotechnology Laboratory, University of Delhi South Campus, New Delhi, grown and maintained on malt extract agar (MEA) containing (g/l) malt extract 20.0, KH₂PO₄ 0.5, MgSO₄.7H₂O 0.5, Ca (NO₃)₂ 4H₂O 0.5, pH 5.4 and at 30 °C[32,33]. This fungal culture has been deposited with International Depository Authority (IDA) at Microbial Type Culture Collection (MTCC- WDCM773), Institute of Microbial Technology (IMTECH), Chandigarh, India (Accession no. MTCC 5722).

Inoculum development. Preparation of pellets in fungal shake flask (250 ml). The 250 ml Erlenmeyer flasks, each containing 50 ml of malt extract broth (MEB) containing (g/l) malt extract 20.0, KH₂PO₄ 0.5, MgSO₄.7H₂O 0.5, Ca (NO₃)₂ 4H₂O 0.5 and pH 5.4, were sterilized at 121 °C (15 psi) for 30 min. Each flask was inoculated with four mycelial discs (8 mm diameter each) taken from the growing edges of 4 day old fungal cultures. The cultures were incubated in an incubator shaker (Innova 44R, New Brunswick Scientific, Enfield, CT, 06082-4444 USA) at 150 rpm and 30 °C for 3 days. The fungal pellets so developed were used as seed inoculum for the scale up of solid state fermentation.

Preparation of fungal pellets in shake flask (2 L). Large scale inoculum production was carried out in 2 L shake flasks, each having 1L of MEB sterilized at 121 °C (15 psi) and inoculated with pellets developed as described above as seed inoculum and incubated at 30 °C for 72 hours and pellets thus developed were used as inoculum for large scale (500 g wheat straw in each tray) development of Biotech Feed.

Solid State Fermentation (SSF) of wheat straw. Laboratory scale. Each enamel tray (30 × 25 × 5 cm) containing 100 g (dry weight) of wheat straw (~5 mm size) moistened with 200 ml (1 : 2) of micronutrient solution was autoclaved at 121 °C (15 psi) for 30 min. The trays were inoculated with fungal pellet inoculum with 1% (w/w) of the substrate on dry weight basis. A total of 6 trays were used in parallel in the lab scale experiment for each incubation period i.e. 5, 10 and 15 days. The trays were incubated at 30 °C for 5, 10 and 15 days. The trays without fungal pellets were used as control. The trays harvested after various fermentation periods were oven dried until it achieved constant weight and processed further for estimation of cell wall composition, proximate content and in vitro digestibility.

Analysis of fungal biomass (ergosterol estimation). Extraction and estimation of ergosterol was carried out with slight modification in the method described by Gattoufi et al. Wheat straw sample (200–250 mg) was macerated with chilled centrifuge tube. The sample was extracted with 5 ml methanol for 2 min using laboratory blender (Model HL 1632/00, Philips India Limited, Gurgaon- 122002, India) and the crude extract was centrifuged at 4500 rpm in a Falcon tube for 5 minutes. The supernatant was then transferred to a 100 ml round bottom flask. The pellet was re-extracted again in 10 ml methanol by shaking at 350 rpm for 10 min followed by a third extraction step. 1.6 g KOH and 4 ml ethanol was then added to the combined supernatants and refluxed for 30 min in a water bath at 80 °C. The saponified solution was filtered by cellulose filters to remove any precipitates and collected in separate funnels. Before extraction, 8 ml distilled water was added and rotated manually for 1 min with three 20 ml portions of petroleum ether (b.p. 35–60 °C). The collected ether phases were evaporated to about 2 ml in a water bath maintained at 30 °C. They were then transferred to small glass tubes and evaporated to complete dryness. The dried sample was re-dissolved in methanol and benzene before estimation using HPLC. The solution was briefly centrifuged, and a volume of 10 μl was injected into a gas chromatograph system having flame ionization detector operating at 282 nm (Waters Corporation 34 Maple Street, Milford, MA 01757 508.478.2000, USA). The column was a reversed-phase, C18 and was protected by a guard column having methanol as mobile phase at a flow rate of 1.5 ml/min. During the entire procedure, samples were protected from direct sunlight.
After washing it in buffer, the samples were post-fixed in 1% OsO4 for 2 hour at 4 °C. The samples were dehydrated in an ascending grade of acetone, critical point dried (CPD) (Critical point dryer, Polaron Quorum Technologies Ltd, Judges House Lewes Road, Laughton Lewes East Sussex BN8 6BN) and mounted on aluminium stubs for SEM. They were sputter-coated (SCD 505 Super Cool Sputter System, Baltec Technology, Raudondvario pl. 148, LT-47175 Kaunas, Lithuania) with colloidal gold and observed under a Leo 435 VP scanning electron microscope (Cambridge, UK) at an operating voltage 15 kV. Images were digitally acquired by using a CCD camera attached to the microscope. For TEM, samples after washing and dehydration in an ascending grade of acetone, samples were infiltrated and embedded in araldite CY 212 (TAAB Laboratories Equipment Ltd 3 Minerva House, Calleva Park Aldermaston, Berks, RG7 8NA, England). Thin sections of grey-silver colour interference (70–80 nm) were cut and mounted onto 300 mesh- copper grids. Sections were stained with alcoholic uranyl acetate and alkaline lead citrate, washed gently with distilled water and observed under a Morgagni 268 D transmission electron microscope (Fei Company, Achtezweg Noord 55651 Gd Eindhoven The Netherlands) at an operating voltage 80 kV. Images were digitally acquired by using a CCD camera (Megaview III, Fei Company) attached to the microscope.

**In vivo digestibility trial.** In vivo animal feeding of fermented feed based diets were carried out after ethical clearance from the Institutional Ethical Committee. Twelve buffalo calves (average body weight 80–85 kg) were purchased locally from Gohana Town, Sonipat, Haryana, India. These calves were divided into three groups and allotted three different treatments/diets. In Treatment I (T1): Wheat straw fed ad libitum with concentrate mixture for maintenance and growth requirement, Treatment II (T2): Biotech Feed fed ad libitum with concentrate requirements for maintenance and growth requirement and Treatment III (T3): Biotech Feed fed ad libitum with concentrate in which 50% grains are reduced. These 50% grains were replaced with Biotech Feed in T3 group, which resulted in 0.34 Mcal less Metabolizable energy/kg Dry Matter in concentrate mixture compared to T1 and T2 groups. Concentration mixture comprised of wheat bran (47.50%) + soybean cake (23.50%) + maize (26%) + mineral mixture (2%) + salt (1%).

**Housing of Experimental Calves.** The calves with identification number on their ears were shifted to a well-ventilated shed having cemented floor. The sheds were equipped to protect the animals from cold and avoid the mixing of feed and residual material of one animal with other. The calves were dewormed before in vivo growth trial.

**Feeding of Calves.** The calves were fed for 90 days for their adaptation with unfermented and fermented wheat straw based diets. Their feeding regime, intake, output, weight gain, and all other parameters were recorded. However, after completion of 90 days, they were further fed for 7 more days and during these 7 days the nutrient digestibility of feed in animals were evaluated. The weighed diets containing either unfermented or fermented wheat straw (Biotech Feed) were fed ad libitum twice a day around 8 A.M. in the morning and 2:30 P.M. in the evening. The concentrate mixture was given according to the specification of National Research Council and on the basis of their growth pattern. The feed offered to particular diet and resident by individual animal were weighed and recorded on daily basis. The animals were fed every day between 6–8 AM in an open area to have exercise and access to running fresh drinking water. The animals were also offered fresh water twice a day.

**Weight gain/weight loss measurement.** Body weight of each animal was recorded at fortnight intervals for three consecutive days at around 8.0 A.M., i.e. before offering the feed and water, and the mean weights were calculated to find out gain or loss in body weight. The calves were weighed on an ‘Avery Make’ bridge type balance (Avery Laboratories Equipment Ltd 3 Minerva House, Calleva Park Aldermaston, Berks, RG7 8NA, England). The samples were collected daily at the time of feeding. All the samples were collected after washing, trimming, and careful to avoid the mixing of feed with remaining feed. Weight loss was calculated by subtracting the dry weight of fermented feed from the weight of unfermented (control) wheat straw28. Oven dried wheat straw (control and fermented ones, diets and feces were ground (1 mm) and sieved (30 mesh) in a moisture free oven at 105 °C for 24 hour or till the weight became constant. The dried samples were collected and taken for dry matter determination. For estimation of nitrogen, an aliquot equal to 1/500 to 1/1000th of the feces voided by the calves were thoroughly mixed with about 1–2 ml of 25% sulphuric acid solution and kept in a wide mouthed glass bottle with a tight lid. At the end of digestibility trial, the bulk samples were ground in a laboratory Willey mill using 1 mm sieve and stored in polythene bags for further biochemical analysis.

**Compositional and proximate analysis.** Loss in dry matter were estimated by weighing the wheat straw before and after fungal fermentation. The weight loss was calculated as changes in its chemical composition, in vitro digestibility and nutrient digestibility in animals.

**Digestibility trial.** Following the 90 days adaptation phase, a 7 days digestion trial was conducted on all the 12 calves used in experimental study. A weighed quantity of diets were offered twice a day at 8.0 A.M. in the morning and 2:30 P.M. in evening. The calves were offered water twice a day. During the digestion trial, samples of unfermented straw, fermented straw (Biotech Feed), green fodder and concentrate mixture were collected daily at the time of feeding. All the samples were collected after thorough mixing of the feed. The residues were also sampled next morning after thoroughly mixing and weighing, 100 g of samples were kept in a hot air oven maintained at 70 °C for 24 hour or till the weight became constant. The dried samples were ground in a laboratory Willey mill (Thomas Scientific, NJ 08085 U.S.A) using 1 mm sieves, mixed well and stored in clean and labeled polythene bags for further biochemical analysis.

**Collection and sampling of feces.** The feces of each calf were collected individually and carefully to avoid the mixing of feces with urine. After 24 hour collection, feces of individual animals were weighed and mixed equally on cemented floor. A weighed quantity of 300–400 g were withdrawn and taken to laboratory for dry matter and biochemical analysis. Out of it, an aliquot equal to 1/50 to 1/100th (the proportion was fixed according to the quantity of feces voided by the individual calves) part of the total feces voided were taken for dry matter determination. For estimation of nitrogen, an aliquot of daily collection equal to 1/500 to 1/1000th of the feces voided by the calves were thoroughly mixed with about 1–2 ml of 25% sulphuric acid solution and kept in a wide mouthed glass bottle with a tight lid. At the end of digestibility trial, the bulk samples were ground in a laboratory Willey mill using 1 mm sieve and stored in polythene bags for further biochemical analysis.

**Statistical Analysis.** The data for in vivo animal feed trial was analyzed using Randomized Block Design (RBD) with three treatments and four replicates as per the procedure suggested by Snedecor and Cochran44. Differences in various treatments were tested by one way analysis of variance (ANOVA) following Bonferroni’s posthoc test (P<0.05) SYSTAST (6.01) software.
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Author contributions

Conceived and designed the experiments: B.S., K.K.J. and R.C.K. Performed the experiments: B.S., K.K.J., R.C.K. and A.K. Analyzed the data: B.S., K.K.J. and R.C.K. Contributed reagents/materials/analysis tools/Animal Facility: R.C.K. and A.K. Wrote the manuscript: B.S., K.K.J. and R.C.K. All authors prepared the manuscript.

Additional information

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