Rapid economical acetic acid Papanicolaou staining procedure versus conventional staining procedure in normal oral mucosa

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Abstract

Background: Papanicolaou (Pap) staining technique is a conventional technique used in cytology but it is time consuming.

Aim: The aim of the study was to determine the efficacy of Rapid Economical Acetic acid Papanicolaou stain (REAP) over conventional staining technique in studying normal oral mucosa.

Materials and Methods: Eighty patients were selected and were grouped based on their habits. Two smears were obtained from each patient and were subjected to both the staining techniques. A total of 160 slides were studied for features such as cellular outline, nuclear outline, nuclear details, cellular differentiation, micronuclei and cellular transparency.

Results: The conventional staining procedure showed 79 cases of optimal cellular staining, 78 cases of optimal nuclear staining, 35 cases of optimal nuclear details, 57 cases of optimal differentiation and 27 cases of optimal transparency and 33 cases of optimal micronuclei. REAP staining showed 75 cases of optimal cellular staining, 64 cases of optimal nuclear staining, 20 cases of optimal nuclear details, 36 cases of optimal differentiation and 28 cases of optimal transparency and 25 cases of optimal micronuclei.

Conclusion: REAP staining effectively reduces the time and the cost factor, but the cytological details are well observed under conventional staining technique in normal patients.

Keywords: Conventional staining, modification, REAP staining

INTRODUCTION

Papanicolaou (Pap) staining technique was first developed by George N. Pap in the year 1943. Pap staining is a commonly used cytological staining technique in screening tests of inflammatory and cancerous lesions of cervix.[1] In the year 1954 and 1960, Pap subsequently introduced modification in the techniques. Various other modifications were introduced aiming to reduce the cost and time involving the procedure.[1] Rapid economical acetic acid Pap (REAP) stain is one such modified staining procedure introduced by Shabnam Izhar et al.[1] The aim of this
The study was conducted in the Department of Oral Pathology and Microbiology, S. R. M Dental College, Chennai. Eighty patients without any oral lesion and inflammation were selected and were divided into four groups. Group 1 (N): patients without habits, Group 2 (S): patients with smoking habit, Group 3 (S + A): patients with smoking and alcohol habit and Group 4 (S + A + P): patients with smoking, alcohol and paan chewing habit. Informed consent was obtained from all the patients before the start of the procedure. Patients were asked to rinse mouth, and the cytological smears were obtained from the buccal mucosa using a wooden spatula with a single stroke. Slides were prepared and wet fixed immediately. The procedure was done twice for each patient to prepare two slides for two different staining techniques such as conventional staining and REAP staining technique.

The staining technique for REAP is:

| Step                                      | Duration |
|-------------------------------------------|----------|
| 1% acetic acid                      | 10 dips  |
| Preheated (60°C) Harris Hematoxylin     | 10 dips  |
| Tap water                              | 10 dips  |
| 1% acetic acid                        | 10 dips  |
| OG-6                                    | 10 dips  |
| 1% acetic acid                        | 10 dips  |
| EA-50                                   | 10 dips  |
| 1% acetic acid                        | 10 dips  |
| Methanol                                | 10 dips  |
| Xylene                                  | 10 dips  |

Cellular outline, nuclear outline, nuclear detail, cellular differentiation, transparency and micronuclei were studied by two independent observers. Based on the visibility of the cellular detail, scores of 0 (suboptimal) and 1 (optimal) were assigned for each feature. The criteria for the scores were:

- Cell outline – Identification of clear, sharp cell outline and absence of fuzziness
- Nuclear outline – Identification of clear, sharp nuclear outline and color intensity
- Nuclear details – Identification of the presence of fragmentation, chromatin network and inclusions in the nuclei
- Differentiation – Identification of cellular color differentiation
- Transparency – Identification of cellular cytoplasm without any particulate
- Micronuclei – Identification of well-defined micronuclei.

RESULTS

A total of 160 slides were studied. The cellular and nuclear details of each patient were studied under conventional and REAP technique. Tables 1-6 represent cellular outline, nuclear staining, nuclear details, cellular differentiation, cellular transparency and micronuclei scores of both techniques.

Under conventional staining technique [Figure 1], the total scores were 79 for optimal cellular outline, 78 for optimal nuclear staining, 35 for optimal nuclear, 57 for optimal differentiation, 27 for optimal transparency and 33 for optimal micronuclei. Scores for REAP staining [Figure 2] were 75 for optimal cellular staining, 64 for optimal nuclear staining, 20 for optimal nuclear details, 36 for optimal differentiation, 28 for optimal transparency and 25 for optimal micronuclei. Significant results were obtained on comparing the two staining techniques without group differentiation in relation to nuclear outline ($P < 0.001$), nuclear detail ($P < 0.013$) and cellular differentiation ($P$ value 0.001). When the groups were divided and studied, in normal patients without habits, conventional staining was better than REAP staining technique in relation to nuclear detail ($P < 0.004$), nuclear differentiation ($P < 0.006$) and cellular transparency ($P < 0.003$). In patients with smoking habits, conventional staining was better than REAP staining technique in relation to cellular transparency ($P < 0.002$). In patients with smoking, alcohol and paan chewing habit, REAP staining was better than conventional staining in relation to cellular differentiation ($P < 0.003$). Apart from the six features that were studied, the study also noticed that the slides stained by either technique showed the presence of inflammatory cells and in few cases presence of bacteria too.

DISCUSSION

Pap staining is by far the most preferred staining method for cytological studies due to its better enhancement of nuclear details and increased cellular transparency. However, it has the limitations of prolonged procedural time and increased costing.

Numerous modifications have been introduced over the years, with the aim of reducing procedural time and cost. Kline, Tao and Sato introduced modification in staining techniques which reduced staining time to 4 min, 5 min and 90 s, respectively.
However, this compromised cytological features.\[^3\]

The ultrafast Papanicolaou stain (UFP) was introduced

| Habit | Cell outline | Staining | \(\chi^2\) | \(P\) |
|-------|--------------|----------|------------|-------|
| N     | Sub-optimal  | 0        | 0          | -     |
|       | Optimal      | 20 (100.0) | 20 (100.0) | 40 (100.0) |
|       | Total        | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S     | Sub-optimal  | 0        | 0          | -     |
|       | Optimal      | 20 (100.0) | 20 (100.0) | 40 (100.0) |
|       | Total        | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A   | Sub-optimal  | 1 (5.0)  | 3 (15.0)   | 4 (10.0)  | - 0.605 |
|       | Optimal      | 19 (95.0) | 17 (85.0)  | 36 (90.0) |
|       | Total        | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A+P | Sub-optimal  | 0        | 2 (10.0)   | 2 (5.0)   | - 0.487 |
|       | Optimal      | 20 (100.0) | 18 (90.0)  | 38 (95.0) |
|       | Total        | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| Total | Sub-optimal  | 1 (1.3)  | 5 (6.3)    | 6 (3.8)   | - 0.210 |
| Optimal | 79 (98.8) | 75 (93.8) | 154 (96.3) |
| Total   | 80 (100.0) | 80 (100.0) | 160 (100.0) |

*Fischer’s exact Chi-square. REAP: Rapid economical acetic acid Papanicolaou

**Table 2: Chi-square test to compare proportions between staining techniques (nuclear outline)**

| Habit | Nuclear outline | Staining | \(\chi^2\) | \(P\) |
|-------|-----------------|----------|------------|-------|
| N     | Sub-optimal     | 1 (5.0)  | 3 (15.0)   | 4 (10.0)  | - 0.605 |
|       | Optimal         | 19 (95.0) | 17 (85.0)  | 36 (90.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S     | Sub-optimal     | 0        | 4 (20.0)   | 4 (10.0)  | - 0.106 |
|       | Optimal         | 20 (100.0) | 16 (80.0)  | 36 (90.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A   | Sub-optimal     | 1 (5.0)  | 5 (25.0)   | 6 (15.0)  | - 0.182 |
|       | Optimal         | 19 (95.0) | 15 (75.0)  | 34 (85.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A+P | Sub-optimal     | 0        | 4 (20.0)   | 4 (10.0)  | - 0.106 |
|       | Optimal         | 20 (100.0) | 16 (80.0)  | 36 (90.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| Total | Sub-optimal     | 2 (2.5)  | 16 (20.0)  | 18 (11.3) | 12.269 <0.001 |
|       | Optimal         | 78 (97.5) | 74 (98.7)  | 152 (98.7) |
|       | Total           | 80 (100.0) | 80 (100.0) | 160 (100.0) |

*Fischer’s exact Chi-square. REAP: Rapid economical acetic acid Papanicolaou

**Table 3: Chi square test to compare proportions between staining techniques (nuclear detail)**

| Habit | Nuclear detail | Staining | \(\chi^2\) | \(P\) |
|-------|-----------------|----------|------------|-------|
| N     | Sub-optimal     | 4 (20.0) | 13 (65.0)  | 17 (42.5) | 8.286 0.004 |
|       | Optimal         | 16 (80.0) | 7 (35.0)  | 23 (57.5) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S     | Sub-optimal     | 13 (65.0) | 17 (85.0)  | 30 (75.0) | 2.133 0.144 |
|       | Optimal         | 7 (35.0)  | 3 (15.0)   | 10 (25.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A   | Sub-optimal     | 17 (85.0) | 17 (85.0)  | 34 (85.0) | - 1.000 |
|       | Optimal         | 3 (15.0)  | 3 (15.0)   | 6 (15.0)  |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| S+A+P | Sub-optimal     | 11 (55.0) | 13 (65.0)  | 24 (60.0) | 0.417 0.519 |
|       | Optimal         | 9 (45.0)  | 7 (35.0)   | 16 (40.0) |
|       | Total           | 20 (100.0) | 20 (100.0) | 40 (100.0) |
| Total | Sub-optimal     | 45 (56.3) | 60 (75.0)  | 105 (65.6) | 6.234 0.013 |
|       | Optimal         | 35 (43.8) | 20 (25.0)  | 55 (34.4) |
|       | Total           | 80 (100.0) | 80 (100.0) | 160 (100.0) |

*Fischer’s exact Chi-square. REAP: Rapid economical acetic acid Papanicolaou
reduction of time to 3 min and cost reduction to 25%.[6]

Bhagat et al. in their study of 102 liquid-based cytology samples that were subjected to Rapid PAP staining technique and the conventional PAP technique reported that the nuclear staining with modified staining was found to be equivalent to conventional Pap technique in 99.9% of cases and the cytoplasmic staining was as good as conventional
technique in 69.6% of cases. The study supported the use of REAP technique as a cost-effective alternative.  

Similar conclusions were drawn by Deshpande and Shabnam Izhar in their studies on the use of REAP technique. Deshpande et al. reported REAP could be used as an alternative to conventional technique in screening programs. The study observed similar quality of nuclear border staining with both conventional and REAP staining technique, but the cell border, cytoplasmic and chromatin staining quality of REAP was slightly inferior to PAP. Excellent slide preservation for 2 years with considerable reduction in time and cost associated with the REAP staining method was also documented by Deshpande et al. Similarly study by Shabnam Izhar reported that the REAP technique is a simple, rapid and economical technique that could be used in places with restricted resources but not for research purpose. In their study of 737 cervical smears, excellent cytomorphological details were observed in 91.6% cases with the use of conventional staining and only in 33.6% cases with REAP staining.

Our study compared the conventional staining and REAP staining technique in normal oral mucosal samples with and without habits. Analysis of total sample data of 80 slides revealed that, except for cellular transparency, greater percentage of optimal reading was observed with conventional staining technique. Statistically significant results were observed with conventional staining techniques in relation to nuclear outline, nuclear detail and cellular differentiation. The procedural time for staining in our series was 3 min due to the use of 1% acetic acid instead of ethanol.

According to the literature, procedures involving acetic acid require less time as it not only helps in nuclear fixation but it also intensifies staining capacity. Acetic acid forms ethyl acetate in combination with ethanol derived from EA-36 and OG-6. This forms complexes with cytoplasm and settle in the cell aiding in preserving the intensity of cytoplasmic staining. This is unlike in the regular conventional procedure where alcohol is used as dehydrant and OG-6 and EA-36stains (alcohol-based stain) diffuse into alcohol during the procedure reducing the cytoplasmic staining intensity.

In our study, although 1% acetic acid was used in the REAP procedure, nuclear outline and details were better observed with conventional PAP staining. Although cellular transparency was higher with the REAP technique, it was not statistically significant.

Asthana and Singh, in their study, compared two stains (conventional Pap and REAP) in 100 smears obtained from 50 normal healthy oral mucosal samples without any habits. The study reported higher optimal cytoplasmic (84%) and nuclear staining (92%) with REAP technique when compared with PAP staining. This was contradictory to our findings where conventional PAP staining was superior or exhibited greater percentage of optimal reading with significant differences in nuclear details (conventional 80%, REAP 35%, $P = 0.004$), differentiation (conventional 90%, REAP 50%, $P = 0.006$) and transparency (conventional 95%, REAP 55%, $P = 0.003$). Although this could be attributed due to the incorporation of samples from participants with habits such as smoking and alcohol which may interfere with the staining reaction, even the normal samples without habits in our study when subjected to both the techniques demonstrated similar outcomes.

When compared to the conventional staining technique, REAP technique was cost effective too as acetic acid was used in the place of ethanol.
CONCLUSION

In this study, cytological features were well observed with conventional staining technique in patients with and without habits. We conclude that though staining time and cost were lower with REAP staining technique were very favorable, conventional staining method was found to be superior to REAP technique.

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Conflicts of interest
There are no conflicts of interest.

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