The application of cyanoacrylate surgical glue on skin suture in rats


Abstract

Purpose: To compare the use of a new cyanoacrylate-based surgical glue and suture with separate points in skin wounds closure.

Methods: Thirty-six rats were subjected to a 4cm dorsal longitudinal incision. Twelve were subjected to simple suture with polyamide 6-0, 12 rats underwent wall synthesis using Dermabond® and 12 was performed cutaneous synthesis with N-2-Butyl-Cyanoacrylate. Twelve of each group was euthanized on the seventh postoperative day, their blood was taken to biochemical tests and a layer of skin and subcutaneous tissue surrounding the surgical scar was randomly divided in two segments, to the submission of tension tests and to histological study.

Results: There were no significant difference between groups (p>0.05). In the soft dermis there was more type I collagen production in group I (p<0.05), group II and III was similar results (p>0.05). In the compact dermis, all 3 groups showed similar results (p>0.05). The biomechanical study was similarity between the glue groups (p>0.05) but the group III proved to be different from the others having a higher resistance (p>0.05).

Conclusion: This glue does not cause any inflammation or kidney and hepatic toxicity. Polyamide sutures are more resistant and the glue should be used alone only in less tension areas.

Key words: Collagen. Wound Healing. Cyanoacrylates. Rats.

DOI: http://dx.doi.org/10.1590/s0102-865020170107

Acta Cir Bras. 2017;32(1):56-64
Introduction

The healing process is a sophisticated mechanism that depends on some factors and they all must be in harmony so the process can develop properly and without complications. It is divided didactically in three phases but they are overlapped and progress continuously, each one of them being critical to the wound healing success.

Cyanocrylates were discovered in 1949 by Ardis and, about ten years later, they had their use documented in surgical procedures. They are liquid monomers that when come in contact with liquids or basic substances become strong glue that is similar to traditional methods of suture when in touch with skin.

During its development, it was observed that despite the strong sealing process, they produced an intense and long-lasting inflammatory reaction. By increasing the size of the chain molecule, these reactions reduced and increased the glue effectiveness.

In tissue healing process, particularly in inflammatory stage, a suture should cause a minimal inflammatory reaction and limited local duration. Nonabsorbable sutures may remain longer in the surgical site causing chronic inflammation and higher chances of infections.

Wound closure techniques researches substantially evolved in the last few years, leading to several studies involving synthetic and absorbable sutures, use of staplers, tapes and other methods. The appearance of surgical glue fits in new methods of wound closure and they have been studied for about four decades.

Surgical glue, besides being an alternative to conventional sutures, have some advantages like technical easiness, decreases surgical time and, since it is not necessary the introduction of a strange substance in the body, decreases wound recovery time. These compounds have been used as complement to conventional sutures. There are studies that show that they are able to create syntheses with no suture in intestines since they do not cause tissue ischemia.

They have unique properties including bacteriostatic effect, hemostatic, it’s biodegradable and biocompatible, except for methyl-cyanoacrylate. They are easily manipulable but show some drawbacks such as lower resistance in high tension areas when compared with conventional sutures and toxicity in some patients, one of the most relevant advantages is the surgical time reduction and the easy handling compared to other methods of synthesis.

2-octylcyanoacrylate became available on the market in German, since then, several studies show that when this glue is used properly, the cosmetic result is similar or above the conventional sutures, with similar rates of infection and dehiscence. 2-octylcyanoacrylate is a glue which creates a layer over the wound keeping the edges together allowing the healing process. It can be used to replace sutures with 5-0 strands or smaller in incisional wounds or lacerations, besides being waterproof.

Although studies demonstrate many benefits, the use of cyanoacrylate glue is not feasible in medical practice due to its high cost. The purpose of this study was to evaluate a new surgical glue composed of N-2-butylcyanoacrylate produced by Universidade Federal do Mato Grosso do Sul (UFMS) chemical laboratory in cooperation agreement with private company under protocol number 23104.001269 / 2010-21 -UFMS by whom this glue is produced with a significantly lower cost than those already on the market.

Methods

This research project was evaluated by an ethics committee in the use of animals.
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(CEUA) of the UFMS and all procedures were performed according to the Animal Experimentation Ethics Committee under protocol number 520/2013, fulfilling the federal law number 11.794, of October 8, 2008 and the decree number 6.689, of July 15, 2009 which regulates the law number 11.794.

Thirty six adult, male rats were used (Rattus norvegicus albinus), weighing between 250g and 350g, coming from UFMS, animal colony.

The animals were identified by tail paint using a combination of three colors, blue, black and green, and separated in three groups of 12 animals each. They were housed in appropriate polypropylene boxes adequate for the species with size of 50x40 cm, lining of wood shavings. They were randomly assigned into three groups:

GI – Containing 12 animals which were shaved and had a 4 cm skin incision in the back followed by cutaneous synthesis using an glue available in the market composed of 2-octylcyanoacrylate (Dermabond);

GII – Containing 12 animals which were shaved and had a 4 cm skin incision in the back followed by cutaneous synthesis using an glue composed of N-2-butylcyanoacrylate;

GIII - Containing 12 animals were shaved and had a 4 cm skin incision in the back followed by cutaneous synthesis using simple suture with polyamide 6-0.

Procedures

Before the beginning of the operative procedures in the surgical technique laboratory of UFMS, the animals were isolated in polypropylene boxes attached to the operating room for a period of 3 hours, aiming to reduce preoperative stress.

For the anesthetic induction the animals were weighted on a precision digital scale and anesthetized with intraperitoneal injection with a solution of 2:1 of ketamine hydrochloride 50mg/ml and xylazine 20mg/ml, respectively, at a dose of (0.1/100g).

Their back hair was cut with scissors followed by mechanical cleaning with soft brush soaked in polyvinyl pyrrolidone iodine 1% and rinse with sodium chloride solution 0.9%.

With animals positioned on operating table in prone position, the back antisepsis was performed with iodine alcohol 2% and sterile surgical cloths were placed isolating the shaved area.

A 4cm incision was made in the skin and subcutaneous tissue of the back in all groups with a 15-scalpel blade, using as references the last rib to caudal margin and spine to dorsal margin, hemostasis was done by compression.

In group I, the defect was restored using 2-octylcyanoacrylate. The group’s II defect was restored using N-2-Butylcyanoacrylate, the edges was approximated and the synthesis made with the surgical glue produced in UFMS, placed on the inner surface of the edges. In group III, the defect was restored using suture with simple stitches with 6-0 polyamide serving as a control group. After the procedures the animals were returned to the same cages used in preoperative, exposed to incandescent light until the anesthetic recovery and maintained with food at will.

After a seven-day observation period (groups GI, GII and GIII) the animals were euthanized by increasing the anesthesia.

Then each rat was immobilized on the operating table, the mechanical suture was removed in group I, in other groups there was no need because the synthesis was made with surgical glue. A segment of skin and subcutaneous tissue of approximately 5x3cm was removed and divided into cranial and caudal parts.
One half was directly taken to a freezer at -20°C to be subjected to the study of rupture strength and the other half was immediately fixed in 10% buffered formalin for the histological analysis.

The samples for rupture strength study were taken frozen for submission to the tensile strength test in the mechanical testing laboratory (National Service of Industrial Learning – SENAI, Teresina – PI, Brazil), using the testing machine DL 20000 (EMIC – Equipment and test systems Ltda, Brazil, Calibrated in 5.22.13 with certificate number 463/13) with manually adjustable gripping claws and electronic data acquisition system, Tesc software version 3.04. The wound line segments were attached to the aluminum claws lined parallel to the synthesis line and the claws were connected to the machine that performed the tension force perpendicular to the synthesis line. The result numbers were provided in kilogram force and converted to gram force. The rupture strength number corresponds to the highest value of the tensile force during the test, in other words, the highest tensile force to break the synthesis line. The traction occurred in a 50mm/min speed and the tensile strength was automatically and constantly measured.

The criteria used for the microscopic study was the quantitative assessment of existing collagen in the skin synthesis observed through the picrosirius red coloration, as well as the classification of maturation (type I and type III). It was used a microscope of Zeiss brand with a x40 magnification and a microcomputer with Proc_image program to quantify the collagen in percentage based on the principles of spectrophotometry. The values reflected a ratio of the amount of collagen in a maximum of 100 points.

For the analysis of biochemical parameters of the cyanoacrylate, it was collected after 7 days 10 ml of blood from each animal by direct puncture into the cava vein. The laboratorial exams were performed in each animal, measuring liver enzymes, aspartate and alanine aminotransferase (ASAT, ALAT), urea and creatinine. For analysis of the inflammatory response was laboratory dosed C-reactive protein (CRP) and serum production of interleukin-6 (IL-6). All tests were performed with quality control of the Brazilian Society of Clinical Analyses (SABC) and American Society of Pathology (SAP).

For the statistical analysis was performed a descriptive analysis of the results found for the serum values of urea, creatinine, ASAT, ALAT, C-reactive protein and interleukin-6 as well as the analysis by Kruskall-Wallis and Brown-Forsythe, in order to compare all 3 groups (Suture with polyamide and synthesis with Dermabond® and N-Butyl 2-Cyanoacrylate).

A descriptive analysis was performed to demonstrate the amount of collagen types I and III found in the experiment. Statistical analysis was performed using 2-way ANOVA test, complemented by Tukey’s test.

For the analysis and comparison of the tensile strength found in each group was applied the Tukey’s test for multiple comparisons.

Results

One of the 36 rats used on the experiment died on the first 24 hours after the procedure, it was therefore excluded from the analysis of the criteria evaluated after 7 days. In the control group animals there were not suture dehiscence observed. In the animals of group I, were observed an amount of 33.3% of dehiscence and the group II presented 41.6% of dehiscence.

The Table 1 shows the results of the descriptive analysis of the parameters studied in each group.
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The normality test indicated a gaussian distribution of data, respecting the homoscedasticity between the groups, reason why variance analysis was chosen. Significant difference was found between the experimental groups (p<0.001), reason why was applied the Tukey’s test for multiple comparisons.

The additional test showed statistical similarities between groups I and II (p>0.05). However, group III proved to be different from the others (p<0.001).

Table 1 - Results found for serum urea (BUN), creatinine, ASAT, ALAT, CPR, IL-6 and rupture strength after 7 days of the cutaneous synthesis. Expressed as mean and standard deviation.

<table>
<thead>
<tr>
<th></th>
<th>Groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Group I</td>
</tr>
<tr>
<td>BUN (mg/dL)</td>
<td>48.3 ± 7.8*</td>
</tr>
<tr>
<td>Creatinine (mg/dL)</td>
<td>0.39 ± 0.03*</td>
</tr>
<tr>
<td>ASAT (U/L)</td>
<td>127.2 ± 35.9*</td>
</tr>
<tr>
<td>ALAT (U/L)</td>
<td>50.67 ± 35.9*</td>
</tr>
<tr>
<td>CPR (mg/dL)</td>
<td>0.20 ± 0.15*</td>
</tr>
<tr>
<td>IL-6 (pg/mL)</td>
<td>1.51 ± 0.03*</td>
</tr>
<tr>
<td>Rupture Strengh (g)**</td>
<td>849.6 ± 403.3*</td>
</tr>
</tbody>
</table>

*p>0.05; **p<0.05

The result of statistical analysis (2-way ANOVA complemented by Tukey’s test) indicates that, in the loose dermis, the group II promoted more formation of type I collagen than other groups (p<0.01), that showed similar results (Table 3).

Table 2 - Average amount of collagen type I and III (in percent) found in the dermis of the tested animals. Expressed as mean and standard deviation.

<table>
<thead>
<tr>
<th>Collagen</th>
<th>GI (%)</th>
<th>GII (%)</th>
<th>Control (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type I*</td>
<td>88.6 ± 3</td>
<td>87.7 ± 2.15</td>
<td>89.5 ± 2.05</td>
</tr>
<tr>
<td>Type III**</td>
<td>11.4 ± 3</td>
<td>12.3 ± 2.15</td>
<td>10.5 ± 2.05</td>
</tr>
</tbody>
</table>

*p>0.01; **p>0.01

Table 3 - Average amount of type I collagen in loose and compact dermis (percent-age) found in the dermis of the animals tested. Expressed as mean and standard deviation.

<table>
<thead>
<tr>
<th>Dermis</th>
<th>GI (%)</th>
<th>GII (%)</th>
<th>Control (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loose</td>
<td>90.2 ± 1.6*</td>
<td>87.4 ± 1.**</td>
<td>87.6 ± 0.9**</td>
</tr>
<tr>
<td>Compact</td>
<td>87.0 ± 2.8**</td>
<td>88.0 ± 3.1**</td>
<td>91.4 ±1.6*</td>
</tr>
</tbody>
</table>

*p<0.01; **p>0.01
In the compact dermis, using the same statistical tests, group III had more type I collagen formation \((p<0.001)\), with the other 2 groups presenting similar results \((p>0.49)\).

When compared only the types of collagen between all the groups, there was no difference when compared type I and III alone, prevailing a greater amount of type I collagen (Figures 1 to 3).

**Discussion**

In this experiment was observed that there was no alteration in the inflammatory tests analyzed in the N-2-butylcyanoacrylate group, showing no systemic inflammatory reaction with the new glue. The values of the tests in the 2-octylcyanoacrylate and control groups with polyamide 6-0 suture also remained at the lower limit. The patterns of CRP and IL-6 were maintained at the device’s minimum reading value, demonstrating no significant difference between the three groups \((p=0.963\) and \(p=0.997\), respectively).

The inflammatory process that may be caused by the toxic products released after the detachment of the cyanoacrylate triggers the release of numerous proteins that can accuse a harmful event when studied and quantified in blood. These markers are pointed as early markers of inflammatory process. The C reactive protein (CRP), an acute phase reactant produced by the liver, has been used in clinical practice with an important scientific support\(^{12}\).

There was no alteration in the renal function when evaluating serum urea levels.
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(p=0.613) and creatinine (p=0.704), showing no renal dysfunction according to the normal parameters reported for this species (45.2 mg/dL–55.0 mg/dL) (0.3 mg/dL – 1 mg/dL), respectively. The kidney is a vital organ with fundamental importance in body homeostasis, receiving approximately 25% of cardiac output. The renal disease is defined as the occurrence of renal morphological lesions or biomechanical abnormality related to the renal function.

The short chain cyanoacrylates stimulate severe inflammatory reaction. The Methyl-2-Cyanoacrylate has necrotizing and pyogenic properties when applied on liver surface, subcutaneous tissue, muscle and bone marrow. Coon et al. demonstrated the development of chronic inflammatory alterations in hearts and kidneys of rats exposed to formaldehyde.

Analyzing the liver enzymes, two animals from group I, six from group II and four from group III had results slightly above the physiological parameters of ASAT (152.4 ± 6.6 mg/dL) and no animal got abnormal ALAT values after 7 days for the species (61.0 ± 2.4 mg/dL), showing that the glue did not cause liver damage.

In wound healing, after 48h of the injury, stands out a fibroplasia phase characterized by an intense fibroblast invasion and multiplication, and healing tissue proteins production and proliferation. The first collagen fibers appear with 4 or 5 days from a fundamental substance produced by fibroblasts (amorphous-mucopolysaccharides) which, in addition to produce these fibers, also guides them. The use of picrosirius red staining was extremely relevant in order to measure the amount and type of collagen fibers, if young or mature, coloring the protein fibers with green when young and red when mature (Type I and III respectively).

After 7 days of experiment was demonstrated that, the three groups presented a higher amount of type I collagen in comparison to type III, in loose and compact dermis, reflecting a wound with greater amount of mature and resistant collagen.

Type I collagen is the most common, synthesized by fibroblasts and is most predominant in bones and tendons. Type III collagen is commonly found in blood vessels, dermis and fascia. Healthy dermis contains about 80% of collagen and 20% of type III.

When compared the three groups only in relation to type I collagen, in this study, the 2-way ANOVA test complemented by Tukey’s test, showed that group I promoted a higher formation of mature or type I collagen than other methods of synthesis (p<0.01), which showed a similar result (p=0.97). In compact dermis using the same statistical methods we had a higher formation of type I collagen in control group (p<0.01). With the other two groups presenting similar results (p>0.49).

It was observed, in this study, a similar rate of short-term dehiscence to literature, showing that this glue is less efficient in high-tension areas than conventional suture, restricting its application to less tension areas.

Despite the many advantages documented by de CA glue, its use remains restricted, due its limited effect in high-tension areas. Using Cavia Porcellus species, Noordzij et al. found significant differences in wound tension immediately after the repair. The authors also concluded that the suture group had a tensile strength 12 times greater than the glue group, but after 7 days, there were no more differences in the tensile strength.

Analyzing animals tensile strength of the cutaneous synthesis, the Tukey’s test showed in both groups I and II significantly weaker wounds after 7 days of observation when compared to the suture group (p<0.001). When comparing both groups submitted to cutaneous synthesis with the surgical CA glue, there was no significant differences between the tensile strength (p>0.05). Previous studies suggest the simultaneous use of porous tapes to help the coaptation of the wound edges and enhance
the synthesis. By analyzing our study and previous authors, it can be concluded that this glue should not be used alone in high-tension areas.

The CA glue use should be limited to non-contaminated or infected wounds, low-tension areas, traumatic or surgical wounds since there is no difficulty in coaptating the wound edges. In fragile skin areas, it has advantages over the suture because it does not cause injury or tissue ischemia.

■ Conclusion

This glue does not cause any inflammation or kidney and hepatic toxicity. Polyamide sutures are more resistant and the glue should be used alone only in less tension areas.

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Conflict of interest: none
Financial source: none

Received: Sep 19, 2016
Review: Nov 18, 2016
Accepted: Dec 20, 2016

1Research performed at Laboratory of Experimental Surgical Technique, Universidade Federal do Mato Grosso do Sul (UFMS), Campo Grande-MS, Brazil. Part of PhD degree thesis, Postgradu-ate Program in Health and Development. Tutor: Ricardo Dutra Aydos.