



貧穉鬱駮糠嗎箕 規床轟靛鑼穉瀦穉駱 粵爛朶韜 驛垢狍濶 農 垢濶咳 秣駮痞

A retrospective study involving 191 patients transplanted from 2007 to 2016 with...
... at department of nephrology, dialysis and transplantation Sahloul Sousse Tunisia...
... consists of the administration of a monoclonal antibody for 67 patients group 1 (G1) and...
... (thymocyte anti-thymocyte globulin or thymoglobulin) for 124 patients group 2 (G2).
... treated with ciclosporin or tacrolimus combined with MMF and corticosteroids or MM...

... transplant patients with mean age of 33.13 ± 13.04 years. The occurrence of episode...
... in patients treated with rATG (21.77% in G2 *versus* 14.92% in G1) but without signif ca...
... ay of occurrence of rejection was shorter in the G1. The uni-varied study showed that th...
... (p=0.005, OR=6.626, IC [1.503-29.20]), urinary tract infections (p=0.020, OR=2.044, CI...
... 38, OR=1.918, CI [1.032-3.564]), CMV infections (p=0.04, OR=2.567, CI [0.996-6.615...
... 035, OR=4.472, CI [0.991-20.186]) are significantly observed with rATG treatment.
... eumopathies (p=0.014, CI [0.034-0.681]) and urinary tract infections (p=0.04, CI [0.27...
... quent with ATG treatment. Neoplastic complications occurred exclusively in G2. W...

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the Student's t test for the comparison of two independent sample means. Alpha risk was judged to be statistically significant from a 5% threshold.

We included 191 kidney recipients. 64.92% of the patients were received as induction therapy a polyclonal antibody and 35.07% received a monoclonal antibody (basiliximub). A high frequency of male sex was observed in both groups, 61.19% and 66.93% respectively with $p=0.427$. There was no significant difference for mean age in both groups, 31.73 ± 13.85 years in G1 versus 33.8 ± 12.57 years in G2, $p=0.279$. The most common initial nephropathy was chronic interstitial nephropathy in both groups, 65.67% and 42.74% respectively with significant difference, $p=0.013$. The mean number of mismatch was higher in the ATG group (3.33 ± 1.60 versus 2.32 ± 1.75) with a significant difference, $p=0.001$. Most patients in both groups received tacrolimus (50.74% in G1 versus 53.22% in G2) with p not significant $p=0.743$. 19.40% of G1 patients received ciclosporin versus 39.51% in G2, $p=0.005$ while treatment with MMF alone was more prescribed in G1 (26.86% in G1 versus 4.03% in G2) with $p=0.001$ (Table 1).

The occurrence of rejection was higher in the group treated with polyclonal antibodies compared with the basiliximub-treated group but without significant difference (21.77% in G2 versus 14.92% in G1), $p=0.253$. The mean time to onset of acute rejection was shorter in the basiliximub group (11.26 ± 21.98 days versus 20.21 ± 44.58 days) with no significant difference $p=0.37$.

Infectious complications were observed particularly in the group treated with polyclonal antibodies with a significant difference for the occurrence of pneumopathies ($p=0.005$), CMV infection ($p=0.045$), urinary tract infections ($p=0.020$), cystitis (0.038) and digestive tract infections ($p=0.035$) (Table 2).

The multivariate analysis revealed that the occurrence of pneumonia ($p=0.014$, IC [0.034-0.681]) and urinary tract infections,

$p=0.04$, IC [0.277-0.969]) were independently associated with treatment with rATG (Table 3).

No patient in group 1 developed neoplasia, while 10 patients in G2 (8.06%) had a neoplastic complication with a significant difference $p=0.017$. There were 3 cases of Kaposi's sarcoma, 2 cases of gastric and cavum lymphoma, 1 tuberkulous adenocarcinoma of the colon, 2 common warts, 2 anal condylomas.

We also evaluated the impact of basiliximub induction versus polyclonal antibody on graft function.

A delayed graft function was observed more frequently in the group treated with rATG 15.32% versus 11.94% but without significant difference $p=0.508$.

Graft loss was observed more frequently in the basiliximub group, 8.95% versus

Pulmonary and digestive infections are independently associated with rATG treatment. This can be explained by the strong immunosuppression induced by polyclonal antibodies. Wang W and