کارگاه‌های آموزشی مرکز اطلاعات علمی

مقاله نویسی علوم انسانی

اصول تنظیم قراردادها

آموزش مهارت های کاربردی در تدوین و چاپ مقاله
Efficacy and Safety of Different Dosages of Praziquantel for the Treatment of Schistosoma Japonicum: A Systematic Review and Meta-Analysis

Damin Cai 1,2; Si Zhang 3; Julong Wu 3; Xun Wang 3; Meng Wang 3; Xiaoling Lu 1; Huiyu Chen 1; Qian Wang 1; Xingming Ma 1,3

1Department of Immunology, School of Basic Medical Sciences, Lanzhou University, Lanzhou, China
2Department of Pathogenic Biology, School of Basic Medical Sciences, Zhejiang Medical College, Hangzhou, China
3Department of Immunology, School of Basic Medical Sciences, Lanzhou University, Lanzhou, China
*Corresponding Author: Xingming Ma, Department of Immunology, School of Basic Medical Sciences, Lanzhou University, Lanzhou, China. Tel: +86-9318289562, E-mail: maxm@lzu.edu.cn

Received: December 11, 2012; Revised: October 28, 2013; Accepted: August 31, 2014

Abstract: Praziquantel, an antischistosomal compound, is used as first-line drug for chemotherapy of Schistosoma japonicum since 1984. In this article, we conducted a systematic review and meta-analysis to evaluate the efficacy and safety of different dosages of praziquantel (PZQ) for treatment of Schistosoma japonicum. The meta-analyses showed that there is no statistically significant difference of the negative rate on the egg using 40 mg/kg compared to 60 mg/kg PZQ for S. japonicum treatment (RR 0.79, 95% CI 0.46 1.35; P < 0.39). The meta-analysis showed that there is no statistically significant difference of the side effects using 30 mg/kg compared with 40 mg/kg (RR 0.97, 95% CI 0.68 1.38; P = 0.87), 40 mg/kg compared with 60 mg/kg (RR 0.97, 95% CI 0.56 1.43; P = 0.63), and 50 mg/kg compared with 60 mg/kg (RR 0.89, 95% CI 0.56 1.42; P = 0.63). According to the results, there is no statistically significant difference in different doses of PZQ for treating S. japonicum.

Keywords: Schistosomiasis Japonicum; Praziquantel; Meta-Analysis

1. Background

Schistosomiasis, an infectious disease caused by parasitic trematodes (schistosomes) dwelling in the host's mesenteric portal system, is a great public health problem in tropical and subtropical regions. The disease causes health and labor loss, and finally a significant reduction in socioeconomic benefits (1, 2). There are five Schistosoma species parasitizing in humans: Schistosoma japonicum, S. mansoni, S. haematobium, S. mekongi, and S. intercalatum. S. japonicum is transmitted by the amphibian snail Oncomelania and causes intestinal and hepatosplenic schistosomiasis in the People's Republic of China, Philippines, and Indonesia; S. mansoni, transmitted by Biomphalaria snails, causes intestinal and hepatic schistosomiasis in Africa, the Arabian peninsula, and South America; S. haematobium, transmitted by Bulinus snails, causes urinary schistosomiasis in Africa and the Arabian peninsula. S. mekongi and S. intercalatum are only of local importance (3-7). In the mid-1980s, the World Health Organization (WHO) recommended schistosomiasis control strategies for humans by focusing on the large-scale population-based and repeated chemotherapy, which is still the key strategy today (8). For schistosomiasis treatment, praziquantel (PZQ) has finally become the first-line medicine. Praziquantel proved to be free of major toxicity, and was well tolerated, highly effective, and easy to administer. Confirmation of results in extended trials may soon permit large-scale treatment (9).

In this article, we conducted a systematic review and meta-analysis to evaluate the efficacy and safety of different dosages of praziquantel (PZQ) for treatment of Schistosoma japonicum.

2. Evidence Acquisition and Analysis

2.1. Search Strategy and Data Sources

We searched the electronic database of PubMed (1966-2012), the Chinese WanFang Literature Database (1992-2012), China National Knowledge Infrastructure (1994-2012.12), and the Chinese Biomedical Literature (1978-2012.12) for all the randomized controlled trials (1992-2012) and the Chinese Biomedical Literature (1978-2012.12) for all the randomized controlled trials evaluating the efficacy and the safety of PZQ with different doses for Schistosoma japonicum treatment. The Review Manager 5.1 software was used to Meta-analysis and using the Revised Jadad scale to access the quality of these studies. The terms and medical subject headings were: "Schistosoma japonicum" or "Praziquantel" or "Meta-Analysis".
(MeSH) used in retrieving citations were “Schistosoma japonicum” (“means the inclusion of all words with the preceding radical), “praziquantel”. The retrieval formula was: Schistosoma japonicum” and praziquantel. The searches were performed mainly in Chinese and English with a limitation to human participants.

2.2. Criteria of Inclusion and Exclusion

After training by a standardized evaluation method, literature were evaluated independently, and screened them in accordance with predetermined. Literature is screened independently by two reviewers. At first, we read the title and summary, and read the full text if the contents were related. All the literature was discussed by two evaluators, if different opinion raised, we discussed together or convinced by our tutor. The inclusion criteria were: (i) independent studies assessing the antischistosomal efficacy of different doses of PZQ, for human schistosomiasis treatment and prevention; (ii) the year of the study conducted or published was reported; (iii) the sample size was reported; (iv) the same drugs’ efficacy evaluation indicators between experimental populations and control populations, i.e. reporting parasitological outcome eggs-positive or eggs-negative by Kato-Katz thick stool smears technique and/or miracidia hatching tests for detecting eggs of S. japonicum, S. mansoni, and S. mekongi or urine filtration for detecting eggs of S. hematobium after approximately 3-4 weeks post-treatment, which was recommended as the standard method for schistosomiasis parasitological diagnosis by WHO in 1980 (8); or reporting emergence or absence of acute schistosomiasis in the trials of assessing some drug’s efficacy in controlling morbidity; (v) the studies were either randomized controlled trials (RCTs) or non-randomized control trials (nRCTs); (vi) reports with raw data, which could be changed into relative risk (RR) and 95% CI RR and 95% CI were reported. Exclusion criteria were: (i) study participants were not human; (ii) without control group; (iii) incomplete information; (iv) duplicate publications; (v) studies described only results without detailed background and method introduction; (vi) reviews. Figure 1 summarizes the studies selection process.

![Flow Diagram Showing the Articles Selection Process for Present Meta-Analyses of the Efficacy Doses of Pzq for Human Schistosomiasis Treatment or Prevention](image)

2.3. Data Extraction and Methodological/Quality Assessment

The extracted information included: first author’s name and year of publication, test sites (i.e. where trials were implemented), time (i.e. when trials were implemented), participants, Schistosoma species, interventions, diagnostics methods, follow-up time, raw dichotomous data of efficacy assessment (NO. of positive/NO. of diagnosed), RRs and their 95% CIs, type of study design (RCT or nRCT), and intervention purposes (prevention or treatment). The quality of included RCTs was assessed by examining whether there is randomization, blinding, and information about follow-up and dropouts/withdrawals of participants according to the guidance of the methodological quality assessment of RCTs in the Cochrane Handbook for Systematic Reviews of Interventions 5.0 and the Jadad scoring criteria (10, 11). The score for quality scale ranges from 0 to 5 points, the higher the score, the higher the quality of trial; and a trial with a Jadad score ≥ 3 has been considered to be of ample quality.
Table 1. Assessment of Methodological Quality of the Included RCTs by Jadad Scoring

| Trial                  | Randomized | Double-Blinded | A Description of Withdrawals or Dropouts | Jadad Score a |
|------------------------|------------|----------------|------------------------------------------|---------------|
| Santos et al. 1979 (9)  | 1          | 2              | 1                                        | 4             |
| Ishizaki et al. 1979 (12)| 1          | 2              | 0                                        | 3             |
| Chen et al. 1985 (13)   | 1          | 1              | 0                                        | 2             |
| Wang et al. 1999 (14)   | 1          | 0              | 0                                        | 1             |
| Wu, 2001 (15)           | 1          | 0              | 0                                        | 1             |
| Li, 2011(16)            | 2          | 0              | 0                                        | 2             |

a Range 0-5 (the higher the Jadad Score is, the higher the quality of the study is).

2.4. Diseases, Interventions and Outcomes

Six articles of Schistosoma japonica were included in this meta-analysis. The participants of experimental groups took different doses of PZQ for treatment. The chemotherapeutic outcome evaluation was parasitological cure, which was defined as eggs-positive or eggs-negative, or emergence or absence of acute schistosomiasis symptoms.

2.5. Data Synthesis and Statistical Analysis

Meta-analyses were conducted in categories of PZQ. RRs based on dichotomous data were set as statistical indicators. Subgroup analyses were conducted according to different design types, different dosages. All the statistical analysis work was performed using the statistical package Review Manager 5 software (Cochrane Collaboration, Oxford, UK) and Stata/SE 11 (Stata. Corporation, Texas, USA) for Egger's publication bias test by LR. The fixed-effects model was used to combine study-specific RRs, when there was no significant heterogeneity among studies. Otherwise, the random-effects model was used.

3. Results

3.1. Studies Selected

Overall, 6 articles met the inclusion criteria and were finally used for this meta-analysis. Figure 1 shows the studies' selection process: The doses of PZQ for treating S. japonicum are 5 mg/kg vs. 10 mg/kg vs. 20 mg/kg (14); 25 mg/kg vs. 40 mg/kg (15); 20 mg/kg vs. 40 mg/kg vs. 60 mg/kg (12); 30 mg/kg vs. 40 mg/kg vs. 50 mg/kg vs. 60 mg/kg (13); 30 mg/kg vs. 40 mg/kg (16); 50 mg/kg vs. 60 mg/kg (9).

3.2. Study Characteristics and Methodological Quality

Studies were conducted in areas that are endemic for Schistosoma japonicum, including Japan, The Philippines, and China (see Table 1). PZQ dosages applied ranged from a single oral dose of 5-60 mg/kg or divided (2, 3) dosages in RCTs-designed studies. For nRCTs about PZQ’s efficacy, there were several types of drug administration i.e. a single oral dose of 40 mg/kg or 60 mg/kg, multiple (2, 3) oral doses of the same concentrations, two doses of 20 mg/kg, and three doses of 20 mg/kg. The follow-up time post treatment differed from studies ranging from 2 days to more than 1 year. Table 1 summarizes the Jadad scores of the included RCTs. Among the 6 RCT-designed studies, all of them have described the randomization method; three study (17) include blinded allocation or outcome measurements, and 1 studies (18) had no description of withdrawals or dropouts. Thus, four of the included RCTs were rated as providing good methodological quality based on a Jadad score of 2-5, and only one study (17) had a Jadad score of 1. The nRCTs without quality assessment were analyzed separately from the RCTs.

3.3. Meta-Analysis

The trials were stratified into sub-groups based on different doses of PZQ. The p value of the test for negative rate on the egg was 0.05, the total pooled ≧ RR, and its 95% CI were calculated by combing all sub-groups. No statistically significant difference among pooled RRs of subgroups about different species was observed (RR 0.79, 95% CI 0.46 1.35, P = 0.39). The trials were stratified into sub-groups based on different doses of PZQ. The p value of the test for negative rate on the egg was ≥ 0.05, the total pooled RR, and its 95% CI were calculated by combing all sub-groups. No statistically significant difference among pooled RRs of subgroups about different species was observed (RR 0.97, 95% CI 0.68 1.38, P = 0.87). The trials were stratified into sub-groups based on different doses of PZQ. The p-value of the test for negative rate on the egg was ≥ 0.05, the total pooled ≥ RR, and its 95% CI were calculated by combing all sub-groups. No statistically significant difference among pooled RRs of subgroups about different species was observed (RR 0.89, 95% CI 0.56 1.42, P = 0.63). The trials were stratified into sub-groups based on different doses of PZQ. The p value of the test for negative rate on the egg was ≥ 0.05, the total pooled RR, and its 95% CI were calculated by combing all sub-groups. No statistically significant difference among pooled RRs of subgroups about different species was observed (RR 0.79, 95% CI 0.46 1.35, P = 0.39) (Figure 2, 3, 4 and 5).
4. Conclusions
Praziquantel was synthesized by Bayer and Merck in Germany in 1972 (19), and introduced for clinical use in the People’s Republic of China since 1978 (19, 20). Today, PZQ is the most frequently used drug for schistosomiasis japonicum treatment.
sis treatment in endemic areas, and regularly used also in large scale programs (21). PZQ exhibits stage-specific functions in killing adult worms (22-24). Our meta-analysis covering a publication period from 1979 to 2011 indicated that PZQ is still effective for S. japonicum with negligible variations. Our meta-analysis showed that different doses of PZQ for treating S. japonicum, which is no significant differences were observed among the 30 mg/kg vs. 40 mg/kg (RR 0.97, 95% CI 0.68 1.38, P = 0.87), 40 mg/kg vs. 60 mg/kg (RR 0.79, 95% CI 0.46 1.35, P = 0.39) or 50 mg/kg vs. 60 mg/kg (RR 0.89, 95% CI 0.56 1.42, P = 0.63). In order to reduce the side effects and the cost of money, we choose the least dose of drug for treating S. japonicum.

Facing the fact that PZQ is still effective for S. japonicum, clinical trials in the future should be specially emphasis on the following aspects: 1) ensure the group good comparability and reduce the generation of selection bias, the random allocation sequence of the report should be detailed and the random program should be hidden; 2) choose the unified international gold standard for diagnosis and severity score; 3) standardize observation time; 4) standardize expression of the data; 5) unify the standards efficacy outcomes and determine the efficacy; 6) Describe the detailed side-effect, ensure the drug is safe.

It is very little quantity of the articles about using different dose of PZQ for the treatment of S. japonicum, so we will the articles like that more and more.

Acknowledgements
We are grateful to Yanping Luo, assistant tutor of the department of Immunology in the school of basic medical sciences in Lanzhou University, and to other colleagues for their kind and very effective cooperation.

Authors’ Contributions
Damin Cai: Designing the study, evaluating the literature, analyzing the data, writing the manuscript. Si Zhang: Evaluating the literature, Julong Wu: Teaching the methods, meta-analysis. Xun Wang: Teaching the methods, meta-analysis. Xiaoling Lu: Screening the literature. Huiyu Chen: Screening the literature. Qian Wang: Screening the literature. Xingming Ma: Designing and supervising the study, evaluating the literature, writing the manuscript.

References
1. Utzinger J, Raso G, Brooker S, De Savigny D, Tanner M, Ornberg N, et al. Schistosomiasis and neglected tropical diseases: towards integrated and sustainable control and a word of caution. Parasitolology. 2009;33(12):1589-74.
2. Hotez PJ, Brindley PJ, Bethony JM, King CH, Pearce EJ, Jacobson J. Helminth infections: the great neglected tropical diseases. J Clin Invest. 2008;118(4):1311-21.
3. Gryseels B, Polman K, Clerinx J, Kestens L. Human schistosomiasis. Lancet. 2006;368(9542):2068-10.
4. Chitsulo L, Engels D, Montresor A, Savili L. The global status of schistosomiasis and its control. Acta Trop. 2000;77(1):41-51.
5. King CH, Dickman K, Tisch DJ. Reassessment of the cost of chronic helmintic infection: a meta-analysis of disability-related outcomes in endemic schistosomiasis. Lancet. 2005;365(9470):156-9.
6. Ross AG, Bartley PB, Sleigh AC, Olds GR, Li Y, Williams GM, et al. Schistosomiasis. N Engl J Med. 2002;346(16):1212-20.
7. McManus DP, Loukas A. Current status of vaccines for schistosomiasis. Clin Microbiol Rev. 2008;21(2):425-42.
8. World Health Organization. Schistosomiasis Unit. The role of chemotherapy in schistosomiasis control. Geneva: WHO Paratic Diseases Programme; 1983.
9. Santos AT, Blas BI, Nosenas JS, Portillo GP, Ortega OM, Hayashi M, et al. Preliminary clinical trials with praziquantel in Schistosoma japonicum infections in the Philippines. Bull World Health Organ. 1979;57(5):785-9.
10. Cochrane Handbook for Systematic Reviews of Interventions Version 4.2.6 [updated September 2006]. Higgins JP, Green S editors. : The Cochrane Collaboration. 2006.
11. Jadad AR, Moore RA, Carroll D, Jenkinson C, Reynolds DJ, Gavaghan DJ, et al. Assessing the quality of reports of randomized clinical trials: is blinding necessary? Control Clin Trials. 1996;17(1):13-132.
12. Ishizaki T, Kamo E, Boehme K. Double-blind studies of tolerance to praziquantel in Japanese patients with Schistosoma japonicum infections. Bull World Health Organ. 1979;57(5):787-91.
13. Chen MGT. Optimum and practical dosification of praziquantel in the treatment of schistosomiasis japonica. Parasitol Parasitic Dis. 1985;2:253-3.
14. Wang YT. Trop Dis Parasitol. 1999;28:198-200.
15. Wu WDT. Trop Dis Parasitol. 2001;30:79-80.
16. Li JM T, Nancharaiah U. 2001;5(3):37.
17. Liu ZG. Effect of intermittent medication with praziquantel for treating schistosomiasis japonica in the endemic season. Chin J Parasit Dis Control. 1997;10(21).
18. Inyang Etoh PC, Ejieze GC, Uche MF, Inyang Etoh EC. Efficacy of a combination of praziquantel and artesunate in the treatment of urinary schistosomiasis in Nigeria. Trans R Soc Trop Med Hyg. 2009;103(1):68-44.
19. Xiao SH. Development of antischistosomal drugs in China, with particular consideration to praziquantel and the artemisinins. Acta Trop. 2000;76(2):315-67.
20. Midzi N, Sangweme J, Zinyowera S, Mapingure MP, Brouwer KC, Kumar N, et al. Efficacy and side effects of praziquantel treatment against Schistosoma haematobium infection among primary school children in Zimbabwe. Trans R Soc Trop Med Hyg. 2008;102(8):759-66.
21. Ma YJ, Guo M, Liu JF. Chemotherapeutic drugs against schistosomiasis japonica. Endemic Dis Bull. 2007;22:68-9.
22. Xiao SH. [Study on prevention and cure of artermether against schistosomiasis]. Chin Jchisto Dis Control. 2003;17(3):310-20.
23. Xiao SH, Yue WJ, Yang YQ, You QJ. Susceptibility of Schistosoma japonicum to different developmental stages to praziquantel. Chin Med J (Engl). 1998;100(9):759-68.
24. Keiser J, Chollet J, Xiao SH, Mei YJ, Jao PY, Utzinger J, et al. Mefloquine–an aminoalcohol with promising antischistosomal properties in mice. PLoS Negl Trop Dis. 2009;3(9).

Cai D et al.
کارگاه‌های آموزشی مرکز اطلاعات علمی

مقاله نویسی علوم انسانی

اصول تنظیم قراردادها

آموزش مهارت های کاربردی در تدوین و چاپ مقاله