Low-dose radiotherapy for COVID-19 pneumonia treatment: case report, procedure, and literature review

Ruben Del Castillo · David Martinez · Gustavo J. Sarria · Luis Pinillos · Bertha Garcia · Luis Castillo · Alicia Carhuactocto · Frank A. Giordano · Gustavo R. Sarria

Abstract

Background The COVID-19 pandemic outbreak has set the emergency services in developing countries on major alert, as the installed response capacities are easily overwhelmed by the constantly increasing high demand. The deficit of intensive care unit beds and ventilators in countries like Peru is forcing practitioners to seek preventive or early interventional strategies to prevent saturating these chronically neglected facilities.

Case presentation A 64-year-old patient is reported after presenting with COVID-19 pneumonia and rapidly progressing to deteriorated ventilatory function. Compassionate treatment with a single 1-Gy dose to the bilateral whole-lung volume was administered, with gradual daily improvement of ventilatory function and decrease in serum inflammatory markers and oxygen support needs, including intubation. No treatment-related toxicity developed. Procedures of transport, disinfection, and treatment planning and delivery are described.

Conclusion Whole-lung low-dose radiotherapy seems to be a promising approach for avoiding or delaying invasive respiratory support. Delivered low doses are far from meeting toxicity ranges. On-going prospective trials will elucidate the effectiveness of this approach.

Keywords COVID-19 · Low-dose radiotherapy · Viral pneumonia · Intensive care · Cytokine storm

Introduction

The coronavirus disease 2019 (COVID-19) pandemic outbreak in late 2019 eventually imposed upon developing countries a major problem for public health systems, due to a lack of installed attention capacity [1]. Examples from European countries such as Spain or Italy, who saw their emergency and hospitalization systems rapidly overwhelmed despite a greater response capacity [2], raised major concern in Latin American countries. In Peru, a major population mobility restriction was introduced on March 15 [3], in order to alleviate the admission rates in a country where, according to official numbers, the overall availability of mechanical ventilators (MV) does not exceed 822 for a total of ~33 million inhabitants [3]. Despite these efforts and as of May 25, the contagion incidence curves seem to still be increasing, with an average of more than 4000 new cases daily since the first official case was reported on March 6 [3, 4].

Currently, in light of the lack of evidence for a specific or effective medical treatment, hospital- and especially intensive care unit (ICU)-based support management have been left as the only options for the expected 5% of advanced cases [5]. According to this and in addition to the previously described situational features, the fragile balance between patients’ admission and attention capacity should be approached from a preventive perspective.

The anti-inflammatory effect of low-dose radiotherapy (LD-RT) was tested for infectious respiratory conditions in former times; however, this practice fell into decline after the advent of medical treatments [6]. Pre-clinical and clinical data regarding toxicity and effectiveness of such treat-
ments have been published, showing an acceptable safety profile [7].

Herein, we present the case of a patient treated at our institution, describe the followed biosecurity and disinfection protocol, and review the available evidence regarding this topic published to date.

**Case description**

We here report on a 64-year-old male patient with a prior history of pulmonary tuberculosis successfully treated in 1975. Ten days prior to admission, he noted malaise, headache, and myalgia, followed by the onset of fever and dry cough 5 days prior to admission. After rapid worsening of symptoms and onset of dyspnea, he presented in the emergency department, where COVID-19 was confirmed by polymerase chain reaction (PCR). Baseline vital signs were: heart rate (HR) 93 bpm, respiratory rate (RR) 22 bpm, temperature (T°) 37.7 °C, blood pressure 133/93 mm Hg, oxygen saturation level (SpO2) 89%.

Oxygen support therapy was started with a non-rebreather mask at 10 L/min output (FiO2 100%) Pa/FiO2 193, with immediate SpO2 improvement to 99%. Chest CT scan revealed a bilateral multifocal ground-glass pattern, with a predominant central and subpleural component. Type I respiratory insufficiency secondary to the viral infection was diagnosed and multidrug treatment including ceftriaxone 2 g intravenous (iv) 24 h, hydroxychloroquine 400 mg per oral (po) twice daily (bid) first day, then 200 mg po bid for 10 days, azithromycin 500 mg po first dose, then 250 mg po per day for 10 days, and enoxaparin 60 mg subcutaneous (sc) per day was started. After 48 h, oxygen support therapy was deescalated to nasal cannula (NC) to 5 L/min, reaching 94% SpO2 and dyspnea pattern improvement. Fever persisted, with T° 39.3 at highest point. Daily laboratory and arterial blood gas (ABG) analyses from therapy installation are summarized in Tables 1 and 2.

During the second day of hospitalization, the dyspnea worsened further and was present at rest. Oxygen therapy with a non-rebreather mask was intensified to 15 L/min, reaching 85% SpO2 and 246 Pa/FiO2. Empiric treatment with tocilizumab 600 mg IV was started and the antibiotic was switched to ceftazidime 2 g iv bid. Due to further deterioration and a high likelihood of fatality of the clini-

| Timepoint | Management onset (3rd and 2nd pre-RT days) | 1st pre-RT day | 24 h post-RT | 2nd day | 3rd day | 4th day | 5th day | 8th day |
|-----------|-----------------------------------|----------------|----------------|--------|--------|--------|--------|--------|
| Hospital area | CIPA | CIPA | CIPA | ICU | ICU | ICU | ICU | CIPA |
| Hemoglobin g/dL (13.5–17.5) | 13.6 | NR | 13.8 | 13.2 | 13.2 | 13.1 | 13.3 | NR | 14.4 |
| Leucocytes/mm³ (4500–11,000) | 4240 | NR | 9190 | 5880 | 5730 | 5290 | 6010 | NR | 5430 |
| Bands (0–400) | 0 | NR | 0 | 0 | 0 | 0 | 0 | NR | 108 |
| Lymphocytes (1300–3500) | 920 | NR | 2417 | 1181.9 | 1048 | 820 | 1220 | NR | 1303 |
| Segmented (1400–6600) | NR | NR | NR | NR | NR | NR | NR | NR |
| Platelets/mm³ (150,000–475,000) | 141,000 | NR | 167,000 | 230,000 | 276,000 | 319,000 | 351,000 | NR | 390,000 |
| D-Dimer mg/L (0.0–0.5) | 0.45 | NR | 0.46 | 0.46 | 1.33 | 1.32 | 1.13 | NR | 0.55 |
| GPT U/L (0.0–41.0) | 61 | NR | 130 | 124 | 117 | NR | NR | NR | 70 |
| AST U/L (0.0–40.0) | 65 | NR | 150 | 99 | 75 | NR | NR | NR | 33 |
| CRP mg/dL (0.0–0.5) | 14.99 | NR | 26.37 | 12.5 | 4.25 | 1.86 | 0.82 | NR | 0.24 |
| Procalcitonin ng/mL (0.0–0.5) | 0.15 | NR | 0.39 | 0.16 | NR | NR | NR | NR |
| Troponin T ng/dL (0.0–0.05) | 0.01 | NR | 0.009 | NR | 0.006 | NR | 0.006 | NR | 0.007 |
| LDH U/L (135.0–225.0) | 393 | NR | 526 | 510 | 429 | 354 | 347 | NR | 260 |
| Ferritin ng/mL (20.0–250.0) | 1817 | NR | 925 | 2418 | 1231.6 | 1034 | NR | 871.2 |
| IL-6 pg/mL (0–7) | 107.1 | NR | 151.1 | NR | 1544 | 1117 | NR | NR | 243.3 |

CRP C-reactive protein, NR not registered, CIPA COVID in-patient area, RT Radiotherapy, ICU Intensive-care unit, GPT glutamate-pyruvate transaminase, AST aspartate aminotransferase, LDH Lactate dehydrogenase, IL-6 Interleukin 6
Table 2  Arterial blood gas analyses (ABG) pre- and post-RT

| Ventilatory device | 3rd pre-RT day | 2nd pre-RT day | 1st pre-RT day | 24h post-RT day | 2nd day | 3rd day | 4th day | 5th day | 8th day |
|--------------------|---------------|---------------|---------------|----------------|---------|---------|---------|---------|--------|
|                    | Baseline      | Non-rebreather mask | HFNC          | HFNC         | HFNC   | HFNC   | HFNC   | HFNC   | Binasal cannula |
| FiO2 (%)           | 21            | 50            | 50            | 7.39          | 7.38    | 7.38    | 7.4    | 7.41    | 30     |
| pH                 | NR            | 7.42          | 7.42          | 7.42          | 7.41    | 7.38    | 7.38    | 7.4     | 7.41    |
| pCO2               | NR            | 38.4          | 41.1          | 40            | 44.8    | 44.9    | 42.5    | 45.3    |
| pO2                | NR            | 193.1         | 73.5          | 85.5          | 88.8    | 90.5    | 85.3    | 80.8    |
| HCO3               | NR            | 22.3          | 24.2          | 24.2          | 25.1    | 25.8    | 25.7    | 25.7    | 28.4    |
| BE                 | NR            | 2.1           | 0.3           | 0.8           | 0.5     | 0.7     | 0.5     | 0.9     | 3.8     |
| SatO2%             | 89            | 93            | 92.5          | 91.2          | 96.6    | 96.9    | 96.9    | NR      | NR      |
| PaO2/FiO2          | NR            | 193           | 104           | 147           | 171     | 222     | 226     | 213     | 269     |
| Lactate            | NR            | 2             | NR            | 3.5           | 2.2     | 2.2     | 1.9     | NR      | 1.9     |

HFNC high-flow nasal cannula, NR not registered

The third day continued with the prior installed management. Clinical and ABG analysis showed RR 25/min, SpO2 91.2%, PO2 73.5 mm Hg, FiO2 0.5, and Pa/FiO2 147. The following protocol was followed regarding biosecurity measures, treatment planning, and delivery: all equipment was precovered with disposable plastic barriers. The patient was transported from the hospitalization unit with a fully isolated transport bed, through a designated transit area at the very end of the regular daily schedule. All involved personnel wore personal protective equipment (PPE), including dedicated garments, goggles, and NIOSH N95 respirators (FFP2 equivalent). A chest planning CT scan was performed, organs at risk (OARs) and whole-lung target volume were contoured. A 1-Gy prescription was opted for and planning with the VMAT technique was performed. Procedure duration included 10 min of patient positioning, 1:20 min CBCT imaging verification, and 1:30 min beam-on time. Taking the same previously described measures, the patient was transported to the intensive care unit (ICU) for continuing medical management. Quaternary ammonium was employed for disinfection and ultraviolet light was additionally used to cover the simulation CT and treatment bunkers.

The patient was kept under observation due to a potential need of intubation and assisted ventilation. Oxygen therapy was continued with a high-flow nasal cannula (HFNC) at 50L/min due to persistent altered ABG values. Medical therapy was intensified with hydroxychloroquine 400 mg bid and azithromycin 500 mg bid; additionally, the enoxaparin dose was intensified (60 mg sc bid).

Three days after RT (sixth hospitalization day), the patient showed improvement in respiratory patterns and a persistent, although ameliorated, cough. HFNC management was de-escalated to a low-flow system with 30% FiO2. ABG parameters showed PaO2 90.5 mm Hg and Pa/FiO2 226.25. Seven days after treatment the patient was discharged from the ICU to continue management in a lower-complexity area. No RT-related toxicity was seen during the process.

Imaging control did not show a clinical correlation: comparing the first CT at admission with the eighth post-RT day, the latter displayed a stronger interstitial inflammatory pattern (Fig. 1).

Table 3  Target coverage and OAR dosimetry

| OAR            | Volume (cc) | Min. dose (cGy) | Max. dose (cGy) | Median dose (cGy) | Dose/volume (cGy) |
|----------------|-------------|-----------------|-----------------|-------------------|-------------------|
| CTV            | 4392.124    | 51.6            | 138.2           | 116.1             | –                  |
| PTV            | 5582.966    | 36.5            | 138.2           | 113               | D95–100           |
|                |             |                 |                 |                   | D50–113.9         |
| Heart          | 615.605     | 59.6            | 123.7           | 89.1              | D2–105.6          |
| Trachea        | 47.747      | 9               | 104.4           | 66.7              | D2–98.2           |
| Spinal cord    | 28.493      | 16.1            | 102.0           | 82.5              | Dmax 100.7        |
| Esophagus      | 28.195      | 7.9             | 109.8           | 78.2              | D2–104            |
| Carina         | 7.857       | 82.2            | 100.6           | 89.6              | Dmax 98.4         |

OAR organ at risk, CTV clinical target volume, PTV planning target volume, Dmax maximum dose
Fig. 1 Baseline and eighth post-RT day imaging. Evident worsening of the inflammatory process can be observed; however, a dissociation between these findings and the good clinical evolution of the patient is to be noted.

Radiotherapy physics and planning considerations

A chest CT scan with 2.5 mm thick slices was acquired for planning purposes, accounting for a total set of 143 images. The hands-up position was determined and non-forced expiration was intended. Delineation of OARs and whole-lung CTV was performed on a contouring system (Monaco v.5.11.02, Elekta AG, Sweden). A circumferential 5-mm and craniocaudal 10-mm PTV expansion was created. The contoured OARs included the heart, esophagus, spinal cord, and trachea. Additionally, the aorta, bronchi, and carina were considered for documentation motives (Table 2). A 1-Gy single dose was prescribed to the PTV (Fig. 2). Planning was performed with the Monte Carlo algorithm, for a 6-MV nominal energy delivered with a 360° arc and 90° collimator position (Elekta Infinity, Elekta AG) for better conformity achievement. Total estimated monitor units (MU) were 692.24. The approximated planning time was 30 min. Equivalent uniform dose (EUD) constraints were employed and priority for heart avoidance was given. Isodose volumes for 1 Gy and 0.5 Gy and dose–volume histograms (DVH) can be seen in Figs. 2 and 3. In addition, the target coverage and OAR dosimetry details can be seen in Table 3.

Literature review and discussion

This first reported case to our knowledge in our country and the region raises the question of the effectiveness of RT to treat COVID-19 pneumonia patients. Due to the lack of contemporary data, it is still difficult to define the safety profile and best treatment timing in terms of prompt or delayed management. Regarding toxicity, the expected secondary effects after whole-lung irradiation are well known by radiation oncologists according to the delivered dose [9, 10]. Doses lower than 1 Gy should not be of major concern for short- or long-term follow-up. Lessons obtained from preclinical and clinical studies, the former performed for the first time over 100 years ago, have already established the basis for this treatment modality [11]. Experiments in cats and mice performed in the 1940s showed a beneficial protective effect in animals exposed to 0.5–1 Gy 24 h after virus inoculation. On the contrary, no greater difference was observed with exposure after 48 h, suggesting a major improvement in cytokine release syndrome (CRS) regulation with early treatment onset [12, 13]. A more recent investigation assessed the impact of a 1-, 2-, and 3-Gy single fraction in a murine model, in order to simulate the impact of radiation exposure in astronauts. After 2-year follow-up, neither of the selected doses caused caspase-3 overexpression nor ceramide species accumulation, which directly induce pneumocyte apoptosis and subsequent fibrosis [14].

The radiobiological rationale for adopting RT as an approach to counteract CRS encompasses a variety of mechanisms. COVID-19 severe acute respiratory syndrome (SARS) pathogenesis is directly related to T helper 1 cell (Th1) activation, which subsequently initiate the dysregulated inflammatory cascade through different factors such as granulocyte colony-stimulating factor (G-CSF), IL-6, IL-7, IL-10, and tumor necrosis factor α (TNF-α) amongst others, and in addition to phagocytic activity, contribute to upregulating the cytokine storm [15–17]. An observed pattern in severe cases points to major damage to tissues expressing high concentrations of angiotensin-converting enzyme 2 (ACE2), mainly type II pneumocytes; therefore, the renin–angiotensin–aldosterone cascade is consequently
added as an extra factor by further activating macrophages and granulocytes and promoting continuous interstitial inflammation [18, 19]. The anti-inflammatory properties of RT include polarization of macrophages to an M2 phenotype (leukocyte adhesion mediator), and downregulation of nitric oxide, TNF-, and TGF-α. Besides, the upregulation of heme oxygenase, IL-10, TNF-β, NFκβ, and apoptosis mechanisms, as well as T-regulatory cell enhancement, have been postulated as co-adjuvant mediatory reactions [20–22]. Moreover, LD-RT might prevent accelerated viral drug-related mutation, which has been recently described due to increased selective pressure, while potentially improving the immune response by means of the previously described mechanisms and enhanced RNA damage compared to antiviral therapy [23, 24].

In the clinical setting, data about nonmalignant pathologies were published during the past century. The comprehensive review published by Calabrese et al. describes a series of publications regarding infectious diseases, including bacterial and interstitial pneumonia [6, 25]. An example is the study published by Oppenheimer in 1943. Satisfactory clinical outcomes were obtained after treating 56 patients with a 0.5-Gy equivalent, doses that were previously reported to satisfactorily treat bacterial lung conditions by Chamberlain [26]. Differences between three subgroups were reported, as patients who were treated from 2–5 days and 6–14 days had full symptom resolution. However, with treatment onset after 14 days, the therapy failed to completely resolve symptoms in 50% of the subjects. No fatalities were recorded [27].

It is also to be remarked that the overall COVID-19 intubation-related mortality rate varies between 30 and 70%, according to specific geographical regions [5, 15, 16]. No major data are available at the moment in terms of lung injury secondary to mechanical ventilation, although this still carries inherent risks [28, 29]. Preventive therapeutic approaches to avoid intubation seem of relevance, as this is associated with elevated mortality per se. However, this premise should not be taken lightly, due to the delicate balance between noninvasive and invasive ventilatory management in these terms. It has been previously demonstrated that noninvasive ventilation and delayed intubation, when indicated, could also yield lethal consequences in SARS management [30, 31].

Regarding irradiation technique selection, a three-arc VMAT plan was preferred over APPA and IMRT-based fields, in order to guarantee whole-lung tissue coverage and OAR (mostly cardiovascular structures) sparing, while simultaneously maintaining the beam-on time as short as possible [32–34]. It should also be noted that positioning and verification times must be accounted for; therefore, expected target coverage, OAR exposure, and image ver-

Fig. 2 1-Gy isodose volume distribution and dose–volume histogram showing a conformal and homogeneous distribution profile on the desired target volume.
ification technique are factors to be included in decision-making. Additional criteria according to available technology and logistic features could be considered. Furthermore, it is important to highlight that the exposure threshold remains controversial for translation into clinical effects. Previous differing publications have described either none or developing of detrimental effects after delivering a 0.5 Gy mean dose to the heart [35, 36]. Additionally, the risk of developing secondary cancers after LD-RT from previously reported benign entity treatments has been estimated to be below 1% [37]. Weighing up all these considerations and due to the rapid deterioration of these patients’ status, prioritizing the management of the acute event, in order to potentially save the patient’s live, is of major relevance.

The compassionate treatment offered to the patient and the possible related response in terms of avoiding invasive maneuvers showed clinical improvement from day 2 after RT. This is also correlated with the observed continuous decrease in inflammatory markers. The initial elevation of serum markers has been described previously, potentially establishing the first downregulator signal and starting point for further inflammatory control; however, this should be further elucidated [38]. Similarly, the observed mild lymphopenia could be derived from the same phenomenon, although the development of COVID-19 has also been reported as a direct cause of this alteration, as for other viral infections [39]. Despite this, the case’s evolution time would indirectly point to a positive effect of RT, due to the usual status worsening of patients after the seventh day. However, no conclusions but only hypothesis-generating premises could be drawn.

Close follow-up, ICU management including continuous ABG, and imaging control are mandatory. The intervention decision should be taken on a multidisciplinary basis and with express consent of the patients, balancing potential benefits and risks, preferably encompassed by a prospective clinical protocol. Ethical concerns are raised as well, due to the rapid deterioration of patients’ status and the need for early decision-making. In desperate need of positive outcomes, errors from selecting inadequate treatment should not be repeated. Recently published evidence from different groups has demonstrated not only the lack of effectiveness but also the risks of employing hydroxychloroquine and azithromycin [40–44].

Despite the previously presented evidence and pathophysiological plausibility, and on the basis of this novel situation, no agent-specific clinical data addressing the possible long-term secondary effects of treating this particular
group of patients are available, nor is it known if a synergy between the COVID-19-induced CRS and RT could impact the incidence of long-term secondary events. This may be taken into consideration when prescribing RT in this setting.

The importance and responsibility of establishing properly designed study protocols to clarify this hypothesis falls to the multidisciplinary team and joint efforts from leading scientific societies. Consensus for patient management and health personnel protection is urgently needed. Results from ongoing trials (NCT04377477, NCT04380818) will elucidate the effectiveness of this therapeutic tool regarding improvement in Pa/FIO₂ parameters, hospital stay, and ICU admission rates. Assessing rates of invasive management requirements after RT as a specific endpoint is also recommended.

Conclusion

Radiotherapy arises as a promising option for COVID-19 pneumonia management. Prospective data from a larger cohort of patients are needed to confirm the safety profile and effectiveness of this approach in this specific group of patients.

Funding  Open Access funding provided by Projekt DEAL.

Conflict of interest  R. Del Castillo, D. Martinez, G.J. Sarria, L. Pinillos, B. Garcia, L. Castillo, and A. Carhuacotocot declare that they have no competing interests. F.A. Giordano reports personal fees from Implacit GmbH, non-financial support from Oncare GmbH, grants and personal fees from NOXXON Pharma AG, grants and personal fees from CARL ZEISS MEDITEC AG, personal fees from Bristol-Myers Squibb, personal fees from Roche Pharma AG, personal fees from MSD Sharp and Dohme GmbH, personal fees from AstraZeneca GmbH, outside the submitted work. In addition, F.A. Giordano has a patent (US 62/435405) pending, not related to the submitted work. G.R. Sarria reports grants and personal fees from Carl Zeiss Meditec AG, not related to the submitted work.

Open Access  This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Castro R, Nin N, Rios F et al (2018) The practice of intensive care in Latin America: a survey of academic intensivists. Crit Care 22(1):39
2. Rudan I (2020) A cascade of causes that led to the COVID-19 tragedy in Italy and in other European Union countries. J Glob Health 10(1):10335
3. Available from: https://pp.pe/politica/gobierno/coronavirus-comando-covid- quedan-185-camas-uci-con-ventiladores-mecanicos-disponibles-de-un-total-de-822-noticia-1262085. Accessed 29 May 2020
4. Torres-Roman JS, Kobiak IC, Valcarcel B et al (2020) The reproductive number R0 of COVID-19 in Peru: An opportunity for effective changes. Travel Med Infect Dis. https://doi.org/10.1016/j.tmaid.2020.101689
5. Thomas-Ruddel D, Winning J, Dickmann P et al (2020) Coronavirus disease 2019 (COVID-19): update for anesthesiologists and intensivists March 2020. Anaesthesist. https://doi.org/10.1007/ s00101-020-00760-3
6. Calabrese EJ, Dhawan G (2013) How radiotherapy was historically used to treat pneumonia: could it be useful today? Yale J Biol Med 86(4):555–570
7. Dhawan Gaurav, Kapoor Rachna, Dhawan Rajiv et al (2020) Low dose radiation therapy as a potential life saving treatment for COVID-19-induced acute respiratory distress syndrome (ARDS). Radiother Oncol. https://doi.org/10.1016/j.radonc.2020.05.002
8. Lara PC, Burgos J, Macias D (2020) Low dose lung radiotherapy for COVID-19 pneumonia. The rationale for a cost-effective anti-inflammatory treatment. Clin Transl Radiat Oncol 23:27–29
9. Bentzen SM, Constine LS, Deasy JO et al (2010) Quantitative analyses of normal tissue effects in the clinic (QUANTEC): an introduction to the scientific issues. Int J Radiat Oncol Biol Phys 76(Suppl):S3–S9
10. Marks LB, Bentzen SM, Deasy JO et al (2010) Radiation dose-volume effects in the lung. Int J Radiat Oncol Biol Phys 76(Suppl):S70–S76
11. Fryer CJ, Fitzpatrick PJ, Rider WD et al (1978) Radiation pneumonitis: experience following a large single dose of radiation. Int J Radiat Oncol Biol Phys 4(11–12):931–936
12. Baylin GJ, Dubin IN, Gobble WG Jr (1946) The effect of roentgen therapy on experimental virus pneumonia; on feline virus pneumonia. Am J Roentgenol Radium Ther 55:473–477
13. Dubin IN, Baylin GJ, Gobble WG Jr (1946) The effect of roentgen therapy on experimental virus pneumonia; on pneumonia produced in white mice by swine influenza virus. Am J Roentgenol Radium Ther 55:478–481
14. Christofidou-Solomidou M, Petrofesa RA, Arguiri E et al (2015) Space radiation-associated lung injury in a murine model. Am J Physiol Lung Cell Mol Physiol 308(5):L416–L428
15. Xiaobo Y, Yuan Y, Jiqian X et al (2020) Clinical course and outcomes of critically ill patients with SARS-CoV-2 pneumonia in Wuhan, China: a single-centered, retrospective, observational study. Lancet Respir Med 8(5):475–481
16. Wang Y, Wang X (2020) COVID-19: a new challenge for human beings. Cell Mol Immunol 17(5):555–557
17. Rothan HA, Byrareddy SN (2020) The epidemiology and pathogenesis of coronavirus disease (COVID-19) outbreak. J Autoimmun 109:102433
18. Vaduganathan M, Vardeny O, Michel T et al (2020) Renin-angiotensin-aldosterone system inhibitors in patients with Covid-19. N Engl J Med 382(17):1653–1659
19. Shi Y, Wang Y, Shao C et al (2020) COVID-19 infection: the perspectives on immune responses. Cell Death Differ 27(5):1451–1454
20. Calabrese EJ, Dhawan G, Kapoor R et al (2019) Radiotherapy treatment of human inflammatory diseases and conditions: optimal dose. Hum Exp Toxicol 38(8):888–898

21. Arenas M, Gil F, Gironella M et al (2008) Time course of anti-inflammatory effect of low-dose radiotherapy: correlation with TGF-beta1 expression. Radiother Oncol 86(3):399–406

22. Frey B, Gaipi US, Sarter K et al (2009) Whole body low dose irradiation improves the course of beginning polyarthritis in human TNF-transgenic mice. Autoimmunity 42(4):346–348

23. Ghadimi-Moghadam A, Haghani M, Bevelacqua J et al (2020) COVID-19 Tragic Pandemic: Concerns over Unintentional “Directed Accelerated Evolution” of Novel Coronavirus (SARS-CoV-2) and Introducing a Modified Treatment Method for ARDS. J Biomed Phys Eng 10(2):241–246

24. Rodel F, Arenas M, Ott OJ et al (2020) Low-dose radiation therapy for COVID-19 pneumopathy: what is the evidence? Strahlenther Onkol. https://doi.org/10.1007/s00066-020-01635-7

25. Rousseau JP, Johnson WM, Harrell GT (1942) The value of roentgen therapy in pneumonia which fails to respond to the sulfonamides. Radiology 38(3):281–289

26. Chamberlain WE (2010) Roentgen therapy with very small doses: experience in lymphadenitis, leukemia, Hodgkin’s disease and certain infections. Acta Radiol 6(1–6):271–280

27. Oppenheimer A (1943) Roentgen therapy of interstitial pneumonia. J Pediatr 23(5):534–538

28. Beitter JR, Malhotra A, Thompson BT (2016) Ventilator-induced Lung Injury. Clin Chest Med 37(4):633–646

29. Gattinoni L, Chiumello D, Rossi S (2020) COVID-19 pneumonia: ARDS or not? Crit Care 24(1):154

30. McEnery T, Gough C, Costello RW (2020) COVID-19: Respiratory support outside the intensive care unit. Lancet Respir Med. https://doi.org/10.1016/s2213-2600(20)30176-4

31. Bellani G, Laffey JG, Pham T et al (2017) Noninvasive ventilation of patients with acute respiratory distress syndrome. Insights from the LUNG SAFE study. Am J Respir Crit Care Med 195(1):67–77

32. Papachristofilou A, Hottinger AL, Weinhold O et al (2019) Heart-sparing volumetric modulated arc therapy for whole lung irradiation. Strahlenther Onkol 195(1):77–82

33. Suzuki G, Ogata T, Aibe N et al (2019) Effective heart-sparing whole lung irradiation using volumetric modulated arc therapy: a case report. J Med Case Rep 13(1):277

34. Kalapurakal JA, Lee B, Bautista J et al (2019) Cardiac-sparing whole lung intensity modulated radiation therapy in children with Wilms tumor: final report on technique and abdominal field matching to maximize normal tissue protection. Pract Radiat Oncol 9(1):e62–e73

35. Tapio S (2016) Pathology and biology of radiation-induced cardiac disease. J Radiat Res 57(5):439–448

36. Tran V, Zablotska LB, Brenner AV et al (2017) Radiation-associated circulatory disease mortality in a pooled analysis of 77,275 patients from the Massachusetts and Canadian tuberculosis fluoroscopy cohorts. Sci Rep 7:44147

37. Trott KR, Kamprad F (2006) Estimation of cancer risks from radiotherapy of benign diseases. Strahlenther Onkol 182(8):431–436

38. Arpin D, Perol D, Blay JY et al (2005) Early variations of circulating interleukin-6 and interleukin-10 levels during thoracic radiotherapy are predictive for radiation pneumonitis. J Clin Oncol 23(34):8748–8756

39. Zhao Q, Meng M, Kumar R et al (2020) Lymphopenia is associated with severe coronavirus disease 2019 (COVID-19) infections: a systemic review and meta-analysis. Int J Infect Dis 96:131–135

40. Tang W, Cao Z, Han M et al (2020) Hydroxychloroquine in patients with mainly mild to moderate coronavirus disease 2019: open label, randomised controlled trial. BMJ. https://doi.org/10.1136/bmj.m1849

41. Rosenberg ES, Dufort EM, Udo T et al (2020) Association of treatment with Hydroxychloroquine or azithromycin with in-hospital mortality in patients with COVID-19 in New York State. JAMA. https://doi.org/10.1001/jama.2020.8630

42. Geleris J, Sun Y, Platt J et al (2020) Observational study of hydroxychloroquine in hospitalized patients with Covid-19. N Engl J Med. https://doi.org/10.1056/NEJMoa2012410

43. Mitja O, Corbacho-Monne M, Ubals M et al (2020) Hydroxychloroquine for early treatment of adults with mild Covid-19: a randomized-controlled trial. Clin Infect Dis. https://doi.org/10.1093/cid/ciaa1009

44. Skipper CP, Pastick KA, Engen NW et al (2020) Hydroxychloroquine in nonhospitalized adults with early Covid-19: a randomized trial. Ann Intern Med. https://doi.org/10.7326/M20-4207