



Impact of Climate Change on Hydrochemical Processes at Two High-Elevation Forested Watersheds in the Southern Appalachians, United States

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Climate change increasingly affects primary productivity and biogeochemical cycles in forest ecosystems at local and global scales. To predict change in vegetation, soil, and hydrologic processes, we applied an integrated biogeochemical model Photosynthesis-EvapoTranspiration and BioGeoChemistry (PnET-BGC) to two high-elevation forested watersheds in the southern Appalachians in the US under representative (or radiative) concentration pathway (RCP)4.5 and RCP8.5 scenarios. We investigated seasonal variability of the changes from current (1986–2015) to future climate scenarios (2071–2100) for important biogeochemical processes/states; identified change points for biogeochemical variables from 1931 to 2100 that indicate potential regime shifts; and compared the climate change impacts of a lower-elevation watershed (WS18) with a higher-elevation watershed (WS27) at the Coweeta Hydrologic Laboratory, North Carolina, United States. We find that gross primary productivity (GPP), net primary productivity (NPP), transpiration, nitrogen mineralization, and streamflow are projected to increase, while soil base saturation, and base cation concentration and ANC of streamwater are projected to decrease at the annual scale but with strong seasonal variability under a changing climate, showing the general trend of acidification of soil and streamwater despite an increase in primary productivity. The predicted changes show distinct contrasts between lower and higher elevations. Climate change is predicted to have larger impact on soil processes at the lower elevation watershed and on vegetation processes at the higher elevation watershed. We also detect five change points of the first principal component of 17 key biogeochemical variables simulated with PnET-BGC between 1931 and 2100, with the last change point projected to occur 20 years earlier under RCP8.5 (2059 at WS18 and WS27) than under RCP4.5 (2079 at WS18 and 2074 at WS27) at both watersheds. The change points occurred earlier at WS18 than at WS27 in the 1980s and 2010s but in the future are projected to occur earlier in WS27 (2074) than WS18 (2079) under RCP4.5, implying that changes in biogeochemical cycles in vegetation, soil, and streams may be accelerating at higher-elevation WS27.

Keywords: PnET-BGC, RCP4.5 and 8.5, Coweeta Basin, change point, seasonal variability

INTRODUCTION

Global climate has changed considerably over the past few decades and these changes will likely continue to accelerate due to increases of greenhouse gas emissions, mainly through burning of fossil fuels and land disturbance (Intergovernmental Panel on Climate Change [IPCC], 2014; Center for Climate and Energy Solutions [C2ES], 2019). Global air temperature between 2081 and 2100 is projected to be 0.3–4.8°C higher than the average between 1986 and 2005 with high confidence (Intergovernmental Panel on Climate Change [IPCC], 2014). Precipitation will likely increase as well (1–4% more per °C increment) with high spatial variability.

Climate change has impacted and will impact the structure, function and productivity of forested ecosystems (McKenney–Easterling et al., 2000; Shugart et al., 2003; McKenney et al., 2007; Albrich et al., 2020) by altering biogeochemical cycles in vegetation, soil, and streams within forests that lead to changes in habitat suitability and species competition. Forests represent 30% of the world land surface and provide a wide variety of ecosystem services, including carbon sequestration, water supply, timber products, wildlife habitats and recreational opportunities (Lindquist et al., 2012; Peters et al., 2013, United Nations, 2020, accessed on 08/01/2020)¹. Climate-driven alteration of biogeochemical cycles will likely impact forests' ecosystem services and associated human use. It is important to predict impact of climate change on forests in order to develop effective mitigation measures to minimize potential adverse consequences to ecosystems and humans.

However, such predictions involve large uncertainties as the response of biogeochemical processes to climate change is complex due to different controlling factors that interact at different spatial and temporal scales (Grimm et al., 2013). Increases in air temperature lead to advancement of spring leaf-out of temperate and boreal trees, and therefore increase annual net primary productivity (NPP) (Zohner et al., 2020). Meanwhile, plant and soil respiration tends to increase with temperature (Turnbull et al., 2001), which results in uncertainty in the direction and magnitude of changes of net ecosystem production. Precipitation and available soil water are critical to forest growth. Changing climate could reduce soil water content due to higher evapotranspiration associated with increase in temperature or changes in plant processes, or it could increase soil water content due to increased precipitation. The change of soil water content and temperature alters soil microbial composition, and biotic and abiotic processes such as decomposition of plant litter and soil organic carbon, nitrification/denitrification, and mineralization of plant nutrients (Stark and Firestone, 1995; Knoop and Vose, 2007; Krishna and Mohan, 2017). Decomposition rates will likely accelerate, therefore, carbon and nutrient (K, Ca, Mg, and P except N) release may become more rapid, especially in the fall and winter (Davidson and Janssens, 2006; Conant et al., 2011). These biogeochemical changes could further impact water availability and quality (Delpla et al., 2009). Floods and droughts will likely become more frequent under climate change (e.g., Wu

et al., 2012, 2014). Water quality may be degraded either because of drought-caused or flood-induced changes in temperature, pH, and concentrations of dissolved oxygen, nutrients, and dissolved organic matter (Stanke et al., 2013).

The response of a dynamic ecosystem such as forests to climate change can exhibit (1) a smooth change, (2) a threshold at particular temperature, precipitation and/or other conditions at which change occurs rapidly, or (3) a transition between two alternate stable states with two thresholds as the environmental driver increases or decreases (Hughes et al., 2013). Detecting the state shift of biogeochemical cycles can be based on rates of change applying change-point analysis (Pedersen et al., 2020). Change-point analysis is a valuable tool to detect abrupt changes in variations or means, and can provide insight on a non-linear response of forest processes to climate change.

In this study, we aim to address the following questions: (1) How does climate change affect vegetation, soil and stream processes and which season shows the largest impact? (2) Do change points of these processes occur over time? Do they correspond to change points of temperature and precipitation? (3) How do the impacts of climate change on forest ecosystem processes differ between lower and higher elevation watersheds?

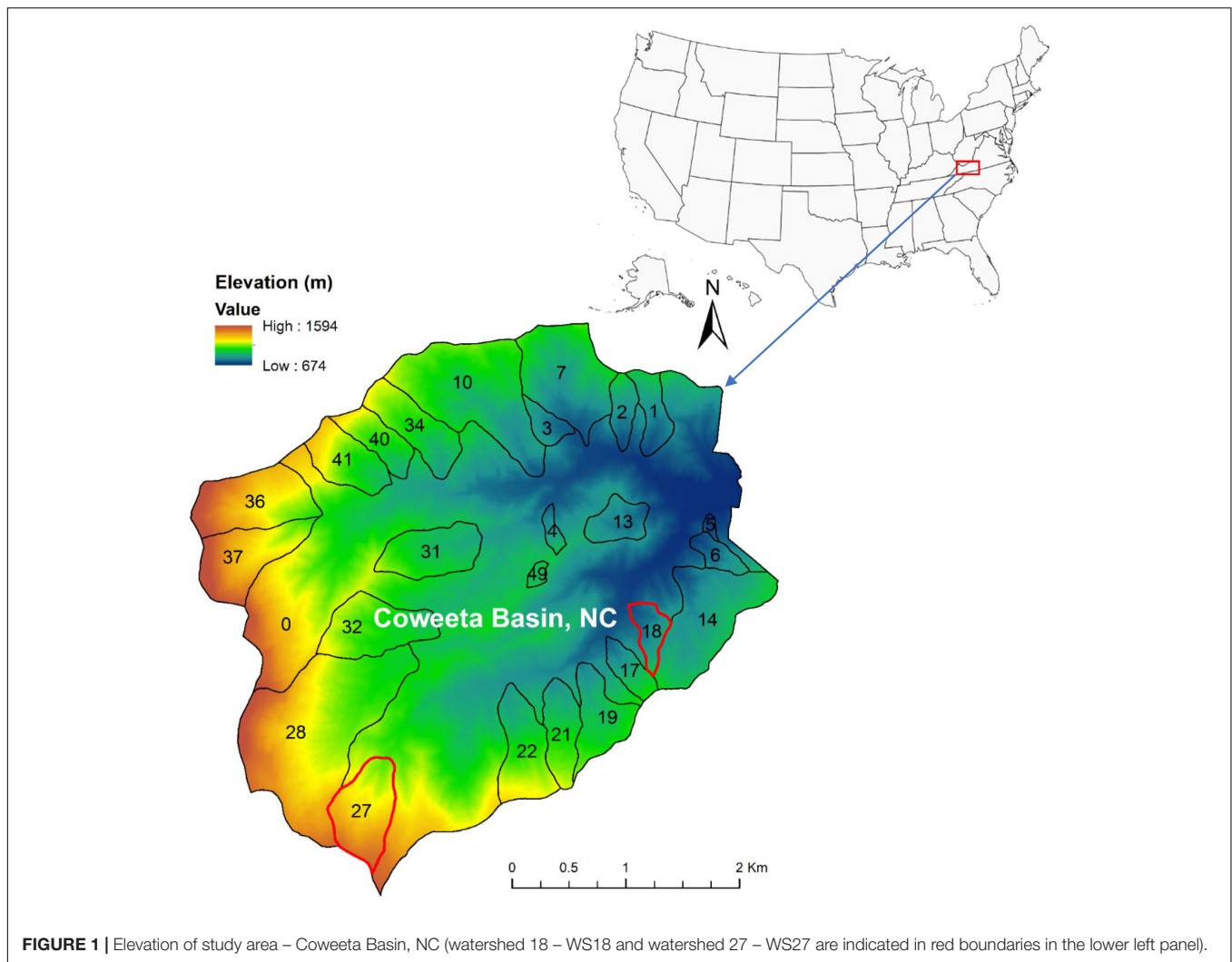
MATERIALS AND METHODS

We applied the PnET-BGC (Photosynthesis-EvapoTranspiration and BioGeoChemistry) model, an integrated and dynamic biogeochemical model (Gbondo–Tugbawa et al., 2001), to evaluate the impact of climate change on coupled vegetation–soil–stream processes at two watersheds at the Coweeta Basin in the southern Appalachian Mountains of North Carolina. We implemented principal component analysis (PCA) to derive the main biogeochemical processes that are important to explain the majority of variance of all ecosystem processes / states and evaluated their seasonal changes. We next derived the change points of ecosystem processes over time.

Study Area

The Coweeta Experimental Forest at the Coweeta Basin was established in 1934 by USDA Forest Service near Otto, North Carolina and was then renamed as Coweeta Hydrologic Laboratory in 1948 (35°03' N, 83°25'W, **Figure 1**). The climate in this region is classified as marine, humid temperate, and features cool summers and mild winters, with less than 5% of annual precipitation occurring as snow (Swift et al., 1988). Two reference watersheds with detailed meteorology, vegetation, soil, and stream chemistry data were selected for this study: lower-elevation watershed 18 (WS18) and higher-elevation watershed 27 (WS27) (**Figure 1**). WS18 is a 12.5 ha watershed drained by Grady Branch, with an elevation range from 726 m to 993 m and a northwest facing slope of 52%. From 1986 and 2015, over half of the precipitation was exported in streamflow (average of 96 cm yr⁻¹ streamflow vs. 182 cm yr⁻¹ total precipitation). The average annual air temperature was 14.0°C with an average minimum of

¹<https://www.un.org/en/observances/forests-and-trees-day>



8.9°C and an average maximum of 19.6°C. WS27 is a 39-ha watershed drained by Hard Luck Creek, and its elevation ranges from 1,061 to 1,454 m with a northeast facing slope of 55%. From 1986 to 2015, over two third of precipitation was lost as streamflow (an average of 163 cm yr⁻¹ streamflow vs. 229 cm yr⁻¹ total precipitation). The average annual air temperature was 10.2°C with an average minimum of 5.9°C and an average maximum of 14.5°C (derived from USDA Forest Service Southern Research Station² accessed on 08/01/2020). Soils are deep sandy loams underlain by folded schist and gneiss. Under the uppermost true and biologically active soils consisting of Ultisols and Inceptisols, there is a porous, friable, and unconsolidated saprolite layer above bedrock, which is believed to be a primary source of base flow and stream geochemistry (Velbel, 1988). Soil depth decreases with elevation with whole basin averaged soil depth around 3 m (Swank and Crossley, 1988). The dominant vegetation at both watersheds is southern mixed deciduous forests with overstory codominance

by oaks (*Quercus*), maples (*Acer*), hickories (*Carya*) and tulip poplar (*Liriodendron tulipifera* L.) and an evergreen understory of rosebay rhododendron (*Rhododendron maximum* L.) and mountain laurel (*Kalmia latifolia* L.) (Day et al., 1988; Elliott and Swank, 2008).

These two watersheds have not been subject to human disturbances since the establishment of the experimental forest in 1934; however, they have experienced a variety of natural disturbances, including chestnut blight in the 1930s (Day and Monk, 1974; Elliott and Swank, 2008), fall cankerworm infestation in WS27 in the 1970s (Swank et al., 1981), extended drought in the 1980s (Elliott and Swank, 1994), and Hurricane Opal in 1995 (Elliott et al., 2002). In addition, the region has experienced increasing air temperature and extreme precipitation events over the years (Laseter et al., 2012).

Model Description, Inputs, and Calibration

PnET-BGC model has been applied, primarily in the northeastern region of the United States, to study forest primary production,

²<https://www.srs.fs.usda.gov/coweeta/tools-and-data/>

water and nitrogen cycling, hydrology, as well as soil and lake/stream acidification in response to acidic deposition, climate change, and disturbances (Gbondo–Tugbawa and Driscoll, 2002; Chen and Driscoll, 2004, 2005a,b; Chen et al., 2004; Zhai et al., 2008; Wu and Driscoll, 2009, 2010; Zhou et al., 2015a; Pourmokhtarian et al., 2016, 2017; Robison and Scanlon, 2018; Valipour et al., 2018). Recently the model has been applied to watersheds in the southeastern and mid and western U.S. (National Smoky Mountain National Park, Zhou et al., 2015b, Niwot Ridge and Loch Vale Watershed, Colorado, and The H. J. Andrews Experimental Forest, Oregon, Dong et al., 2019). With the work presented herein, Coweeta Basin represents the southernmost U.S. application of PnET-BGC. This model includes two major modules: (1) PnET, a submodel that simulates water, carbon, and nitrogen cycling through vegetation, soil and drainage waters, and (2) BGC, a module that includes vegetation and organic matter interactions of abiotic soil processes, solution speciation, and surface water process involving other major elements (Gbondo–Tugbawa et al., 2001). The main state variables that are simulated include primary productivity, evapotranspiration, streamflow soil exchangeable cations, carbon and nitrogen mineralization, litter biomass, concentrations of sulfate, nitrate, potassium, calcium, magnesium, sodium, and acid neutralizing capacity (ANC) in stream at a monthly time step. We applied the model version that accounts for the impact of CO₂ on primary productivity based on the results from the Duke University's Free-Air Carbon Dioxide Enrichment (FACE) project (Schlesinger et al., 2006) where gross primary productivity (GPP) increases with atmospheric CO₂ concentration until 600 ppm. Increasing atmospheric CO₂ has two confounding effects that are depicted in this model: an increase in maximum photosynthetic rate ($A_{\max}$) and a reduction in stomatal conductance (g_s). Stomatal conductance and photosynthesis are coupled (Jarvis and Davies, 1998; Ollinger et al., 2002, 2009). Stomatal conductance changes in proportion to the difference between atmospheric CO₂ (C_a) and internal CO₂ (C_i) across the boundary of stomata. C_i is estimated from C_i/C_a (Ollinger et al., 2009, $C_i/C_a \sim 0.7$ in Lavergne et al., 2019). The stomatal conductance and maximum photosynthetic rate are used to calculate CO₂ assimilation. Additionally, water use efficiency (WUE) is a function of CO₂ assimilation and vapor pressure deficit (VPD), adjusted by C_i/C_a and increases in $A_{\max}$ ($\Delta A_{\max}$). When atmospheric CO₂ increases, $A_{\max}$ will increase and g_s will decrease, which results in an increase of WUE. Plant transpiration is constrained by both $\Delta A_{\max}$ and adjusted conductance (Δg_s) when C_a is high and is calculated from GPP and WUE. We further modified the PnET-BGC model by adding a base flow component in order to obtain more accurate simulations of streamflow. The previous PnET-BGC model would simulate zero streamflow if no precipitation occurred in that month, which is not consistent with the observed streamflow pattern (such as December 2006 at WS18, and October 2000 at WS27). Based on observed non-zero streamflow when monthly precipitation was zero between 1936 and present, we added base flows of 0.6 cm/mo at WS18 and 0.9 cm/mo at WS27 to the streamflow originally simulated by the PnET-BGC.

In order to run the PnET-BGC, meteorology and atmospheric deposition data of major elements are required as inputs (Supplementary Tables 1, 2).

Meteorological Data

Meteorological inputs include monthly maximum and minimum air temperature ($T_{\max}$ and $T_{\min}$, °C), photosynthetic active radiation (PAR) ($\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$), and precipitation ($\text{cm}\cdot\text{mo}^{-1}$). The daily maximum and minimum temperature were available from 1985 (WS18) /1992 (WS27) to 2015 (see footnote 2 last accessed on 05/28/2020). The longer monthly time series of temperature was available at Climate Station 1 (CS01) in Coweeta (1935–2015). We derived regression models based on monthly temperature measured at CS01 and at WS18/27 (WS18: 1985 to 2015, $R^2 = 0.90$ for maximum temperature: $T_{\max-\text{WS18}} = 0.9968 \times T_{\max-\text{WS01}} - 0.6957$, and $R^2 = 0.91$ for minimum temperature: $T_{\min-\text{WS18}} = 1.0029 \times T_{\min-\text{WS01}} + 1.8471$; WS27: 1992–2015, $R^2 = 0.96$ for maximum temperature: $T_{\max-\text{WS27}} = 1.0137 \times T_{\max-\text{WS01}} - 6.1392$, and $R^2 = 0.96$ for minimum temperature: $T_{\min-\text{WS27}} = 1.0091 \times T_{\min-\text{WS01}} - 0.7467$). After deriving climate data from 1935 to 1985 or to 1992 for WS18 and WS27 based on the regression models, the monthly means of maximum and minimum temperature from 1935 to 1985 or to 1992 were used as temperature inputs before 1935 for each watershed. The monthly averages from 2016 to 2100 in the base climate scenario were derived using monthly mean temperatures from 2006 to 2015 (Hansen et al., 2006).

Monthly precipitation was available from August 1936 to 2015 at WS18 and April 1958 to 2015 at WS27. We derived the average total precipitation values for each month and applied these values to before 1936 and to the future (2016–2100) in the base climate scenario.

Measured PAR was available only for a limited period: May 2010 to December 2011 at WS18 and 27. To obtain a longer-term series of PAR, we applied the simulated solar radiation from 1980 to 2010 based on the Daily Surface Weather and Climatological Summaries (DayMET), we converted solar radiation to PAR using Eq. 1. (Both et al., 2002):

$$48.3 \text{ W m}^{-2} \text{ (solar radiation)} = 100 \mu\text{mol m}^{-2} \text{ s}^{-1} \text{ (PAR)} \quad (\text{Eq. 1})$$

The derived monthly averages of PAR between 1980 and 2010 were used as inputs for the years before 1980 and after 2011 under the base climate scenario.

The climate change scenarios were statistically downscaled from four global atmosphere-ocean general circulation models (AOGCMs: CCSM4, HadGEM2, MIROC5, and MRI-CGCM3) with two representative concentration pathways (RCPs, RCP4.5, and RCP8.5). A station-based asynchronous regional regression model was based on the long-term, and daily observed climate data collected from Coweeta WS18 (1985–2015) and WS27 (1992–2015) (Hayhoe et al., 2004, 2007, 2008; Pourmokhtarian et al., 2016). The spatial resolution of the AOGCMs (~ 100 km) is too coarse for application to WS18 or 27 because small

watersheds in complex mountainous terrain are strongly affected by local weather patterns (Pourmokhtarian et al., 2012). The station-based technique generally returns higher precipitation projection than the bias correction-spatial disaggregation grid-based downscaling technique, another common downscaling approach (Pourmokhtarian, 2013). As climate change scenarios can contribute considerable uncertainty in model projections (Huber et al., 2021), it is important to consider multiple climate change scenarios. The RCP8.5 represents the most aggressive greenhouse gas (GHG) emission scenario. By 2100, the atmospheric CO₂ concentration is projected to be 1,370 ppm (Moss et al., 2010). RCP4.5 represents a scenario with some mitigation plans, and atmospheric CO₂ concentration is projected to reach 580 ppm in 2100. To simplify the study and follow atmospheric CO₂ trends between 1970 and 2016 (a linear increase from 330 to 403 ppm), we applied a linear extrapolation to derive annual predictions of atmospheric CO₂ concentrations from 2016 to 2100 as done in other studies (Kienast, 1991), based on the target CO₂ concentrations in 2100 in RCP4.5 and RCP8.5 and CO₂ concentration of 403 ppm in 2016. The predicted temperature and precipitation changes in all four seasons under climate change are similar between WS18 and 27 under each RCP scenario, except that the increase of precipitation under RCP8.5 at WS27 is predicted to be smaller than that at WS18, especially in spring and summer. The RCP8.5 scenario shows higher temperature and precipitation increase than RCP4.5 and temperature is projected to increase from past (1936–1965) to current (1986–2015) and then to future conditions (2071–2100), while precipitation initially decreased from past to current then increase in the future.

We further applied paired *t*-tests to evaluate whether annual temperature or precipitation in the future (2071–2100) under changing climate are significant different from the present (1986–2015). We first checked the normality and temporal correlation (acf function in R) for annual time series between 1986 and 2015 and between 2071 and 2100, and found that data followed normal distribution and there did not exist temporal correlation, therefore justifying the use of paired *t*-tests.

Wet and Dry Depositions

The wet and dry deposition of major elements (Na⁺, Mg²⁺, K⁺, Ca²⁺, NH₄⁺, Cl[−], NO₃[−], and SO₄^{2−}) were obtained from National Atmospheric Deposition Program (NADP, accessed on 08/01/2020)³ and The Clear Air Status and Trend Network (CASTNET, accessed on 08/01/2020)⁴ respectively. The closest NADP site to the Coweeta Basin is NC25 (35°03′36.1″ N, 83°25′50.7″ W, located within the Coweeta Basin⁵, accessed on 08/01/2020) and the wet deposition record is from July 1978 to 2015. For the dry deposition, the CASTNET COW137 station (35°03′36.1″ N, 83°25′50.7″ W) located adjacent to the NADP station, was used with the data available from November 1987 to 2015. Atmospheric deposition data before 1978/1987 was reconstructed from national emission record

from 1930 to 1978, and the relationship between emission and atmospheric deposition of SO₂ and NO_x between 1978 and 2015 (United States Environmental Protection Agency [USEPA], 2000; Driscoll et al., 2001; Chen et al., 2004; Chen and Driscoll, 2004). From 1850 (pre-industrial) to 1930 when the emission data were not available, we assume a 50% increase of SO₂ and NO_x emission during this period and a linear increment was applied to reconstruct sulfur and nitrogen emissions and wet deposition from 1850 to 1930. After wet deposition data were fully constructed, the monthly average dry-to-wet-deposition-ratios derived from observed CASTNET and NADP data were used to reconstruct missing dry deposition data for the period outside the period of observations from the CASNET station. In addition to S and N (SO₄^{2−} and NO₃[−]), other chemical constituents include NH₄⁺, Ca²⁺, Mg²⁺, K⁺, Na⁺, and Cl[−]. For these elements, re-construction of their deposition data before and after the observed NADP record period (i.e., before 1978 and after 2016), 5-year averages of monthly values were applied respectively (i.e., monthly deposition before 1978 was calculated using observations between 1979 and 1983 while we used averages from 2011 to 2015 for the time after 2016). For those elements that are not measured through NADP (dissolved organic carbon – DOC, Al³⁺, F[−], PO₄^{3−}, and Si), we used the input data in reference to previous work from the northeastern U.S.⁶ (accessed on 08/01/2020).

Vegetation Data

The PnET-BGC model previously considered four vegetation types: Northern Hardwoods, Spruce – Fir, Red Oak-Red Maple, and Red Pine. In this study, we constructed and applied Southern Hardwoods vegetation type based on the parameters of the Northern Hardwoods vegetation type. Particularly, we adjusted the optimum temperature for photosynthesis (28°C at Watershed 18 and 23°C at Watershed 27), growing-degree-days at which foliar production begins and ends (400 and 1500 respectively at Watershed 18, 350 and 1400 respectively at Watershed 27), and fraction of precipitation intercepted and evaporated (0.20 at Watershed 18, and 0.11 at Watershed 27). The adjustments are based on previously measured data of NPP and continuous measured data of streamflow.

Calibration

To calibrate the model, we applied the data provided by the Coweeta Hydrologic Laboratory, including streamflow, stream concentrations of Na⁺, Mg²⁺, K⁺, Ca²⁺, Cl[−], NH₄⁺, NO₃[−], and SO₄^{2−}. Other data used to calibrate different modules of the model included NPP, net soil mineralization rates, and soil base saturation available in the literature (Day and Monk, 1977; Knoepp and Swank, 1998; Knoepp et al., 2016; **Supplementary Table 3**).

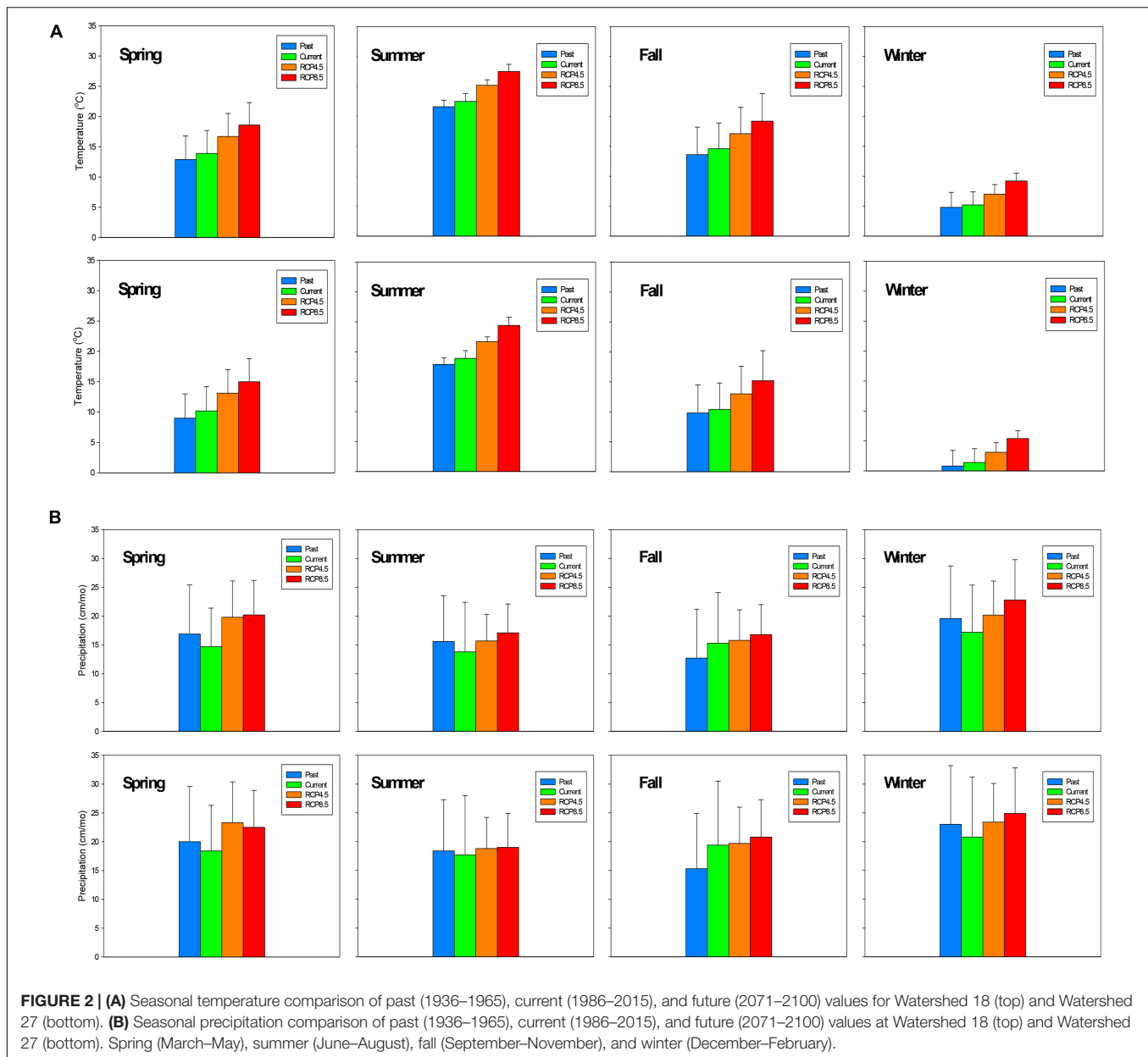
To quantify how well model simulations matched the site observations, we applied normalized mean error (NME) and normalized mean absolute error (NMAE, Janssen and Heuberger, 1995). The NME (Eq. 2) provides the average prediction bias and NMAE (Eq. 3) indicates the absolute error between predicted

³<http://nadp.slh.wisc.edu/>

⁴<https://www.epa.gov/castnet>

⁵<http://nadp.slh.wisc.edu/data/ntn/plots/ntntrends.html?siteID=NC25>

⁶<https://ctdrisco.expressions.syr.edu/pnet-bgc-model/>



and observed values. NME and NMAE closer to 0 indicates a better model fit.

$$NME = \frac{\bar{p} - \bar{o}}{\bar{o}} \quad (\text{Eq. 2})$$

$$NMAE = \frac{\sum_{t=1}^n (|p_t - o_t|)}{n\bar{o}} \quad (\text{Eq. 3})$$

Where p is model prediction and $\bar{p}$ is the mean of predictions; o is observation and $\bar{o}$ is the mean of observations, and n is the number of observations.

Principal Component Analysis

To reduce the data dimension of the many coupled vegetation-soil-stream processes studied, we applied principal component analyses based on 17 vegetation-hydrology-soil-stream variables simulated from 1931 to 2100 under both RCP4.5 and 8.5 climate scenarios with the PCA analysis in base R (<https://www.r-project.org/>, last accessed on 08/01/2020). The 17 variables considered are GPP, NPP, total litter mass, streamflow, evapotranspiration, soil base saturation, soil Al:Ca, net nitrogen mineralization rate, gross nitrogen mineralization rate, gross nitrogen immobilization rate, nitrogen uptake rate, concentrations of NO_3^- , SO_4^{2-} , Ca^{2+} , Mg^{2+} , and K^+ , and the acid neutralizing capacity (ANC) of streamwater ($\mu\text{eq/L}$). Note our application of PCA is for data

TABLE 1A | Seasonal temperature and precipitation in the past, current, and future (RCP4.5 and RCP8.5) conditions at WS18 and WS27.

		Spr	Sum	Fall	Win	Mean	Differ. ²	Var. ³
<i>Climate¹</i>		Temperature (°C)						
WS18	Past	12.9	21.6	13.6	4.8	13.2		
	Current	13.9	22.5	14.6	5.2	14.1	0.8*	
	RCP4.5	16.7	25.2	17.1	7.0	16.5	2.5^	0.5
	RCP8.5	18.6	27.5	19.2	9.2	18.6	4.6^	0.4
WS27	Past	9.0	17.9	9.8	0.8	9.4		
	Current	10.2	18.9	10.4	1.4	10.2	0.9*	
	RCP4.5	13.1	21.7	13.0	3.1	12.7	2.5^	0.5
	RCP8.5	15.0	24.4	15.2	5.4	15.0	4.8^	0.6
Precipitation (cm/mo)								
WS18	Past	16.9	15.6	12.7	19.6	16.2		
	Current	14.7	13.8	15.3	17.2	15.3	−1.0*	
	RCP4.5	19.8	15.7	15.8	20.2	17.9	2.6^	1.9
	RCP8.5	20.2	17.1	16.8	22.8	19.2	4.0^	2.0
WS27	Past	20.0	18.4	15.3	23.0	19.2		
	Current	18.4	17.7	19.4	20.8	19.1	−0.1*	
	RCP4.5	23.3	18.8	19.7	23.4	21.3	2.2^	2.0
	RCP8.5	22.5	19.0	20.8	24.9	21.8	2.7^	1.6

¹Past climate refers to the period between 1936 and 1965, current climate between 1986 and 2015, and future climate between 2071 and 2100 from four GCMs under either RCP4.5 or RCP8.5 scenario.

²"Differ" refers to mean of difference of climate between scenarios, either current climate minus past climate (*) or future climate minus current climate (^).

³Seasonal variability – variance of future climate minus current climate. Seasons: Spr – March to May, Sum – June to August, Fall – September to November, and Win – December to February.

dimension reduction and identification of important watershed process variables. Due to the exploratory nature of this analysis, normality of the data is not a strict requirement (Vaughan and Ormerod, 2005; Jolliffe and Cadima, 2016).

Seasonal Analysis

We evaluated the changes of the ecological processes/states under changing climate for spring (March to May), summer (June to August), fall (September to November), and winter (December to February). We tested whether the 17 variables simulated from the PnET-BGC model are significantly different under changing future climate (2071–2100) compared to the present (1986–2015) in each season using paired *t*-tests. We first checked the normality and temporal correlation (acf function in R) for annual time series (one seasonal average for each season of each year) between 1986 and 2015 and between 2071 and 2100, and found that data followed normal distribution and there did not exist temporal correlation, therefore justifying use of paired *t*-tests.

We focused on the variables that are important in Principal Component 1 and 2 and these include transpiration, GPP, NPP, soil base saturation, N mineralization, base cation (particularly potassium and calcium) concentrations, and ANC in streamwater.

Change-Point Detection

Identifying a change point in the time series is the first step in identifying a potential driver of change, and, therefore a

TABLE 1B | Statistical test of temperature and precipitation between current and changing climate scenarios.

Temperature								
RCP4.5 vs. Current					RCP8.5 vs. Current			
	Spr	Sum	Fall	Win	Spr	Sum	Fall	Win
WS18	***	***	***	***	***	***	***	***
WS27	***	***	***	***	***	***	***	***
Precipitation								
WS18	***	**	—	—	***	***	*	—
WS27	***	**	—	—	***	***	—	—

*Indicates $0.01 < p < 0.05$.

**Indicates $0.001 < p < 0.01$.

***Indicates $p < 0.001$.

— Indicates no significant difference has been detected.

Color green means increasing from current to the future. In each season from 1986 to 2000 and from 2071 to 2100, monthly climate data were averaged first, then the 30 data points in each of the current and changing climate scenarios were checked for temporal autocorrelation ("acf" function in R) and normality (Shapiro test). As the checks show lack of temporal autocorrelation and the data met normality requirement, *t*-test was implemented to test whether the seasonal climate variables under current and changing climate scenarios were statistically significant different or not.

TABLE 2 | Comparison of observed and simulated streamflow (unit: cm/month) and stream chemistry variables (unit: $\mu\text{mol/L}$) at WS18 and WS27.

	Observed		Simulated		Model performance	
	mean	SD*	mean	SD	NME	NMAE
WS18						
Na ⁺	40.4	2.0	41.6	2.2	0.02	0.08
Mg ²⁺	12.4	0.6	11.5	0.7	0.01	0.09
K ⁺	11.3	0.9	10.7	0.4	0.01	0.08
Ca ²⁺	15.7	1.0	16.4	0.8	0.002	0.08
Cl [−]	14.6	0.6	16.0	2.2	0.000	0.12
NO ₃ [−] _N	0.8	0.5	0.3	0.2	0.65	0.74
SO ₄ ^{2−} _S	4.7	1.2	6.0	0.5	0.30	0.38
ANC	83.7	5.7	80.6	5.2	0.04	0.08
Streamflow	8.2	2.5	9.0	2.3	0.09	0.15
WS27						
Na ⁺	22.1	1.6	22.3	2.0	0.02	0.09
Mg ²⁺	8.4	0.6	8.7	0.7	0.03	0.08
K ⁺	5.8	0.7	5.6	0.1	0.04	0.09
Ca ²⁺	8.9	0.9	8.4	0.5	0.06	0.08
Cl [−]	13.2	1.4	13.5	1.8	0.02	0.08
NO ₃ [−] _N	2.3	1.1	1.1	0.9	0.50	0.69
SO ₄ ^{2−} _S	11.8	1.6	11.3	1.1	0.05	0.13
ANC**	23.6	2.5	25.3	4.0	0.07	0.13
Streamflow	14.1	3.3	14.0	3.0	0.003	0.06

The water chemistry data ranged from 1972 to 2014 (ANC from 1973 to 2014), streamflow ranged from 1936 to 2015 at WS18, and from 1947 to 2015 at WS27.

*SD denotes standard deviation. NME is normalized mean error, NMAE is normalized mean absolute error. **ANC denotes acidic neutralizing capacity (Driscoll et al., 1994).

mechanism for a potential regime shift (Anderson et al., 2009), even though the existence of an abrupt change point does not necessarily lead to instability, hysteresis or a regime shift. To

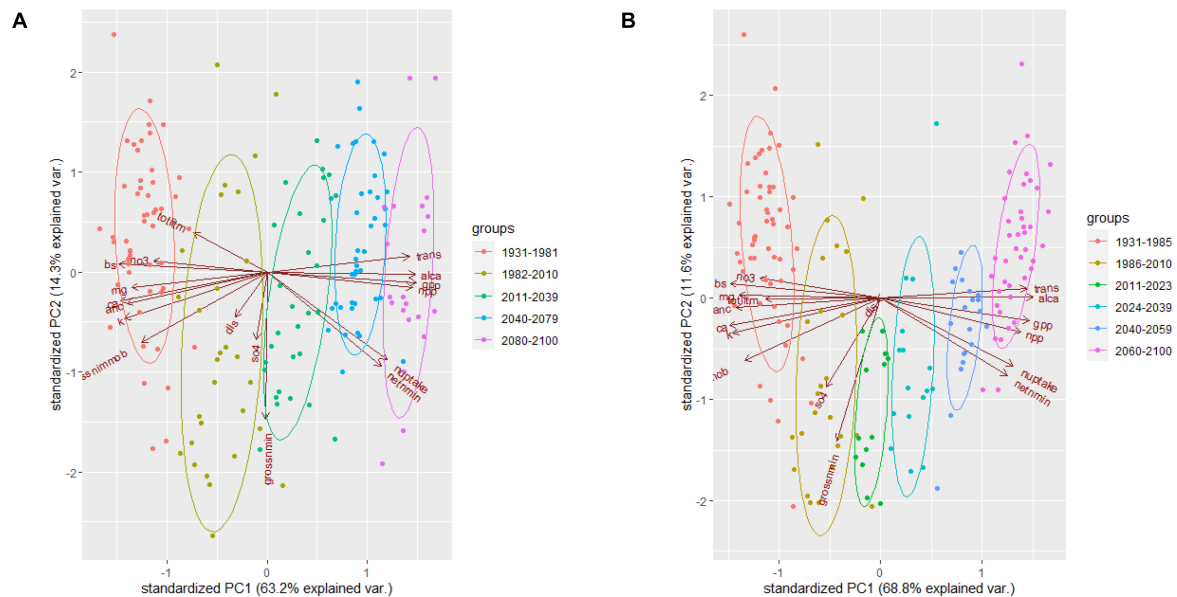


FIGURE 3 | Principal component analysis (PCA) for the 17 variables at WS18 under RCP4.5 (A) and RCP8.5 (B). *trans*, transpiration; *alca*, ratio of Al and Ca in soil; *gpp*, gross primary productivity; *npp*, net primary productivity; *nuptake*, nitrogen uptake by vegetation; *netnmin*, net mineralization rate of nitrogen; *grossnmin*, gross mineralization rate of nitrogen; *SO4*, sulfate concentration in stream; *dis*, stream discharge; *grossnimmob*, gross immobilization rate of nitrogen; *k*, potassium concentration in stream; *anc*, acid neutralizing capacity; *ca*, calcium concentration in stream; *mg*, magnesium concentration in stream; *bs*, base saturation in soil; *no3*, nitrate concentration in stream; *totlitr*, total litter mass.

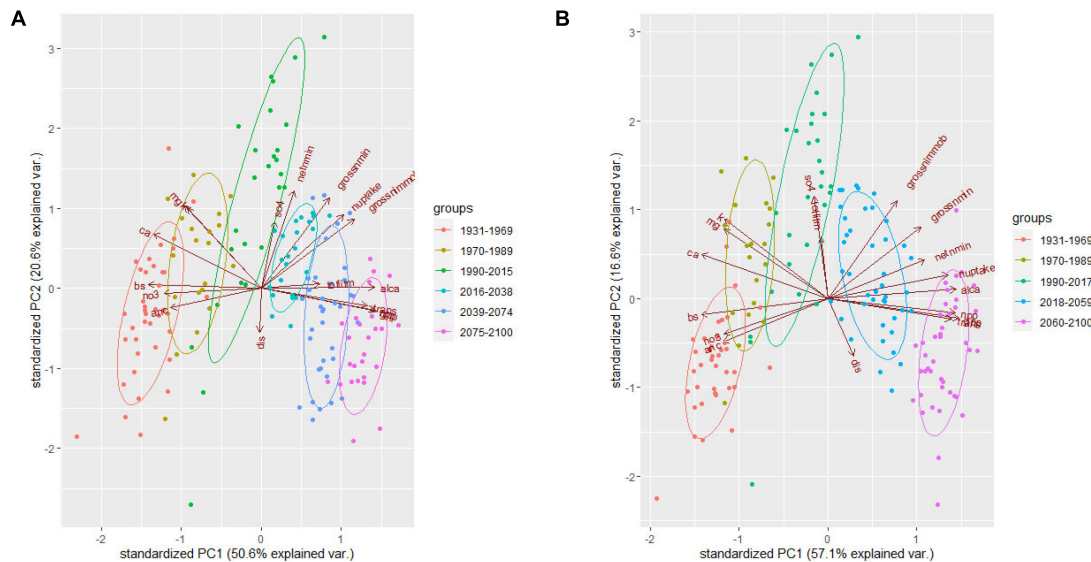


FIGURE 4 | Principal component analysis (PCA) for the 17 variables at WS27 under RCP4.5 (A) and RCP8.5 (B). *trans*, transpiration; *alca*, ratio of Al and Ca in soil; *gpp*, gross primary productivity; *npp*, net primary productivity; *nuptake*, nitrogen uptake by vegetation; *netnmin*, net mineralization rate of nitrogen; *grossnmin*, gross mineralization rate of nitrogen; *SO4*, sulfate concentration in stream; *dis*, stream discharge; *grossnimmob*, gross immobilization rate of nitrogen; *k*, potassium concentration in stream; *anc*, acid neutralizing capacity; *ca*, calcium concentration in stream; *mg*, magnesium concentration in stream; *bs*, base saturation in soil; *no3*, nitrate concentration in stream; *totlitr*, total litter mass.

further investigate the impact of climate change on watershed processes, we conducted change-point analyses in the time series (1931–2100) of PCA score means of the first principal components (PC1s) that explained the largest variance of the

variables (Anderson et al., 2009), to identify the years when PC1s exhibited or are projected to exhibit significant shifts, using the R-package of “changepoint” (Killick and Eckley, 2014). We have selected the binary segmentation (“BinSeg” in R) method.

We checked for the normality between each change point (i.e., years) identified and they all met the normal distribution requirement (on annual basis). The same change point detection methods were used for the climate data between 1931 and 2100 with climate data after 2016 being the predictions for each RCP scenario (RCP4.5 and 8.5). For both PC1s and climate data (1931–2100) change-point detections, we implemented the program to allow the number of change points to vary from three to ten. However, our analysis consistently revealed only five change points.

RESULTS

Our research revealed three key findings. (1) The impacts of climate change on the forest ecosystem at Coweeta vary by season. There is not a particular season during which impacts are most prevalent, but rather the seasonal response depends on the biogeochemical processes considered. (2) Five change points were found over the interval 1930–2100 for PC1s, which explained 50–60% of the variance of the key

biogeochemical processes/states, driven primarily by temperature or precipitation. (3) Vegetation seems to be impacted by climate change more at the higher-elevation WS27 than lower-elevation WS18, while soil and stream processes seem to be more impacted at WS18.

Climate Change

For the two studied watersheds in the Coweeta Basin, observed air temperature increased from the past (1936–1965) to the current condition (1986–2015) and is projected to continue to increase between 2.5 and 4.8°C from current conditions to the future (2071–2100) (**Figure 2** and **Table 1A**). WS27 shows a slightly greater temperature increase than WS18 from current to the future conditions, particularly under RCP8.5 (+4.6°C at WS18 vs. +4.8°C at WS27). Under RCP4.5, temperature increase in spring is slightly larger than in summer followed by fall and winter (increase of 2–3°C except winter with an increase of < 2°C). Under RCP 8.5 summer has the largest temperature increase followed by spring, fall, and winter (increase of 4.5–5.5°C

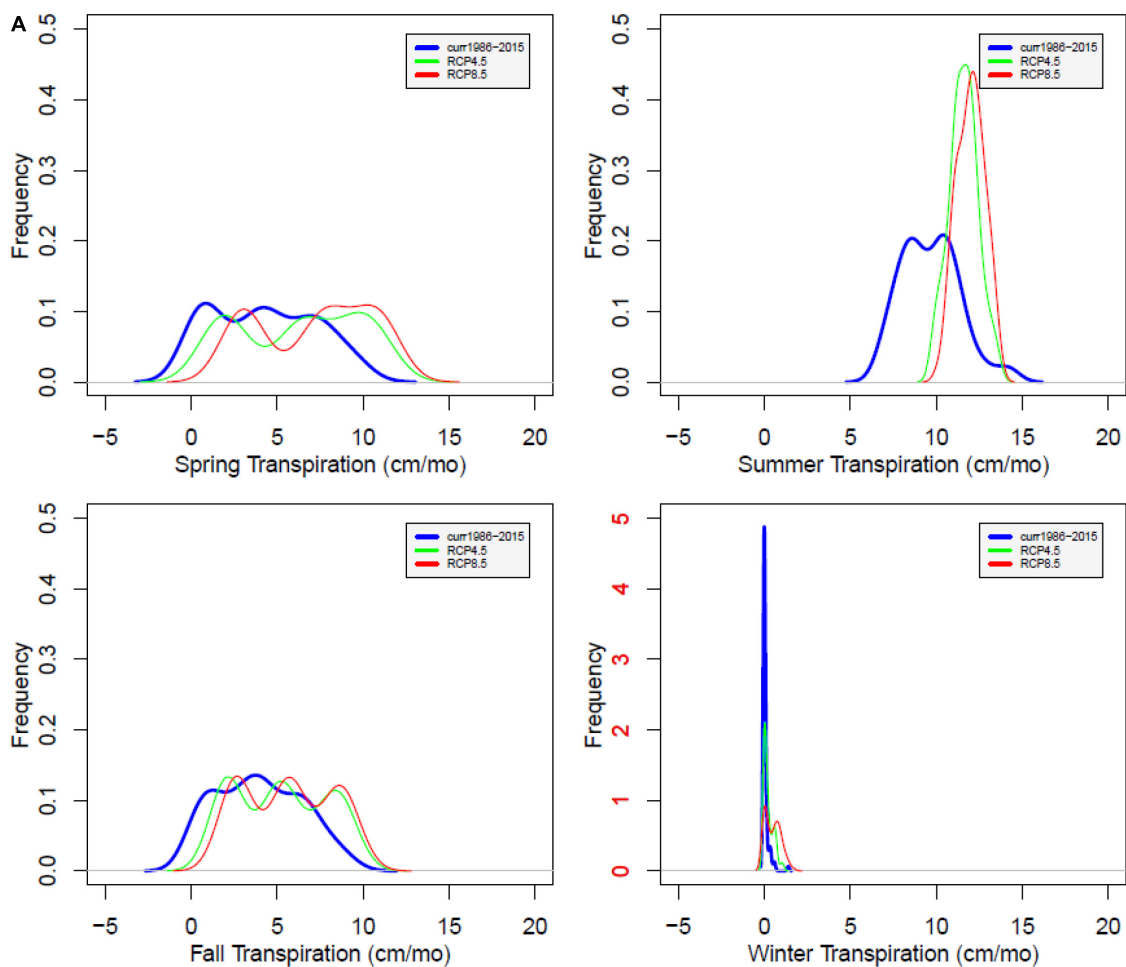


FIGURE 5 | (Continued)

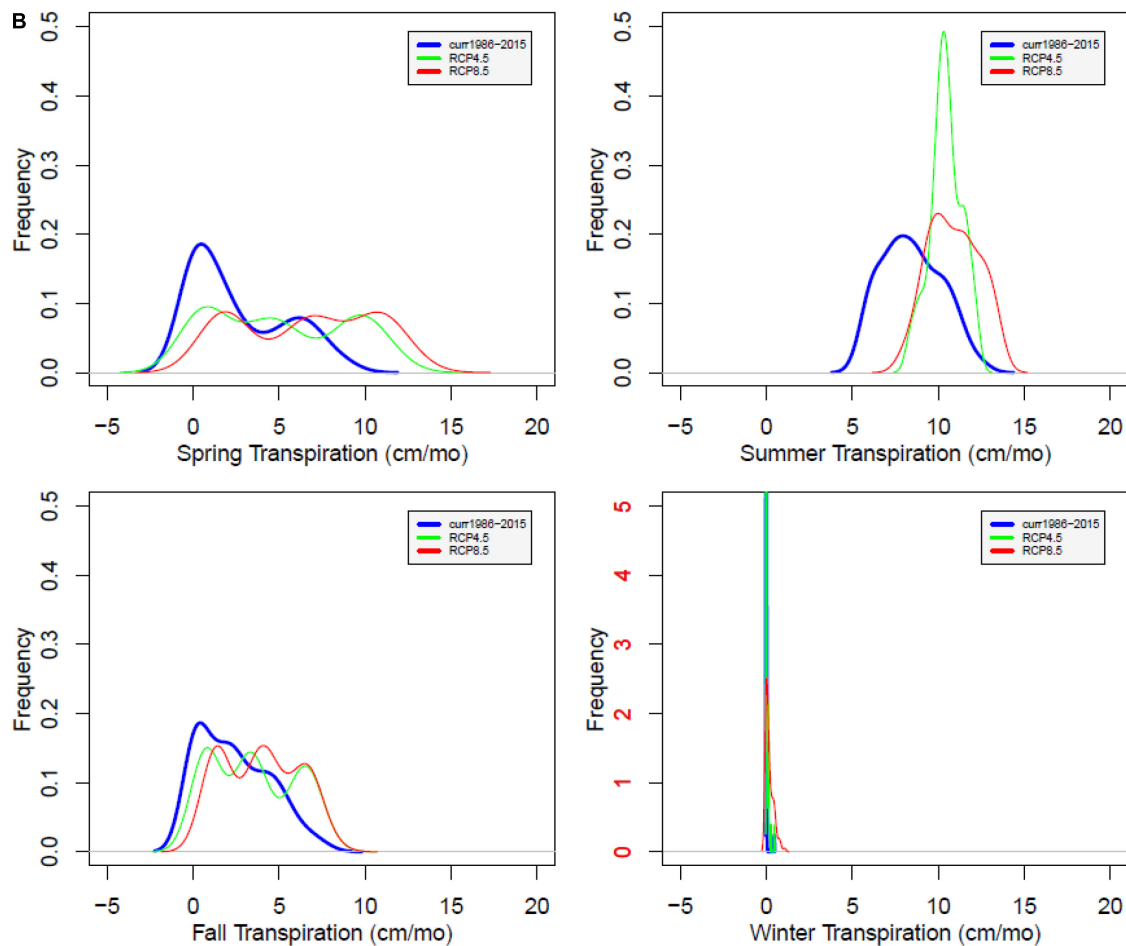


FIGURE 5 | Probability distribution of simulated transpiration at WS18 (A) and WS27 (B) under future climate compared to values under current climate by season. Current (blue), RCP4.5 (green), and RCP8.5 (red). The red color of the labels in winter shows different range of y-axis compared to the other seasons.

except winter with an increase of 4°C) (Table 1A and Figure 2A).

Observed precipitation decreased from past to current conditions during all the seasons except for fall. In contrast, future precipitation is projected to increase ranging from 2.2 to 4.0 cm/mo in all seasons in the future compared to current conditions with the fall showing the smallest increment (Table 1). Both air temperature and precipitation projections for Coweeta follow a similar pattern of regional climate projections for the southeast U.S. (Intergovernmental Panel on Climate Change [IPCC], 2014). Noticeably, the temperature and precipitation projections for each season under changing climate scenario (2071–2100) are statistically significantly different from those under current climate scenario (1986–2015) (Table 1B).

Model Performance

The PnET-BGC mostly effectively simulated monthly and annual vegetation-soil-stream processes at both WS18 and WS27 based on the stream chemistry data from 1970s to

2014 and streamflow data from 1930s/1940 to 2014, with NME ranging from −0.06 to 0.09 (Table 2). However, NO_3^- was underestimated at both WS18 and WS27 and SO_4^{2-} was overestimated at WS18. The model also underestimated the variance of NO_3^- and SO_4^{2-} . Measured concentrations of SO_4^{2-} at WS18 showed an increase between 1980 and 1990s that are seemingly inconsistent with long-term decreases in atmospheric deposition of S during the same period after implementation of the Clean Air Act (1970) and Clean Air Act Amendments (1990) (USEPA⁷ accessed on 08/01/2020). However, this pattern may reflect desorption of soil SO_4^{2-} to streamwater that was absorbed during prior decades of elevated atmospheric S deposition (Rice et al., 2014). The nitrogen cycle is more complex than other element cycles and it strongly linked to processes of vegetation and microbial activities, and therefore, PnET-BGC simulation of stream NO_3^- involves larger uncertainties than other solutes (Fakhraei et al., 2016; Shao et al., 2020). In addition, contrary to a decrease in both observed nitrogen

⁷<https://www.epa.gov/clean-air-act-overview/evolution-clean-air-act>

deposition and simulated nitrate leaching since the 1970s, both watersheds show an unexplainable increasing trend of observed nitrate concentration in streams, which the model does not reproduce.

In addition to the long-term observed streamflow and stream chemistry concentrations, our model simulations were consistent with other published observed data (**Supplementary Table 3**), including NPP, soil base saturation, and net nitrogen mineralization rate. Model simulated average aboveground NPPs at WS18 and WS27 between 1971 and 1980 are close to published data from Day and Monk (1977) (8,056 vs. 7,965 kg/ha-yr). For net nitrogen mineralization rate, the 10-year averages (1971–1980) are 2.5 and 3.1 mg N/kg soil-mo at WS18 and WS27, which are within the range reported by Knoepp and Swank (1998) based on the samples collected along an elevation gradient (<1.2 – 3.8 mg N/kg soil-mo), and consistent with the pattern that the highest values occur at elevations over 1,000 m. WS18 at the lower elevation has higher soil base saturation (BS) of 24.9% than higher elevation WS27 with a lower BS of 4.0% with simulated data from 2001 to 2010 (**Supplementary Table 3**). This simulated range is close to the data range obtained from USDA Forest Services in 2008 soil survey from 9.2 to 19.7%.

Principal Component Analysis

The PCA analysis indicated that the first three principal components explained ~ 85% of variance of the 17 variables analyzed (**Supplementary Tables 4, 5** and **Figures 3, 4**). Particularly, PC1 explained 63.2 and 68.8% of total variances under RCP4.5 and RCP8.5 respectively at WS 18. PC1s explained 50.6% and 57.1% of the variances under RCP4.5 and RCP8.5, respectively, at WS 27. Based on the eigenvectors, the first principal components (PC1s) represent the contrast between primary production and transpiration vs. soil and stream ANC, and the second principal components (PC2s) depict nitrogen mineralization rate. The third principal components (PC3s) represent the concentrations of sulfate in stream at WS18 under RCP4.5 and at WS27 under both RCP4.5 and RCP8.5, while the PC3 at WS18 under RCP8.5 represents streamflow.

Seasonal Variability

Based on the PCA analysis, the evaluation of the impacts of climate change on watersheds of Coweeta focused on the processes that are important in PC1s and PC2s during different seasons. Particularly we focus on seasonal changes of transpiration, NPP, and stream ANC in the main text. The figures for other process variables are described in Appendix Figures

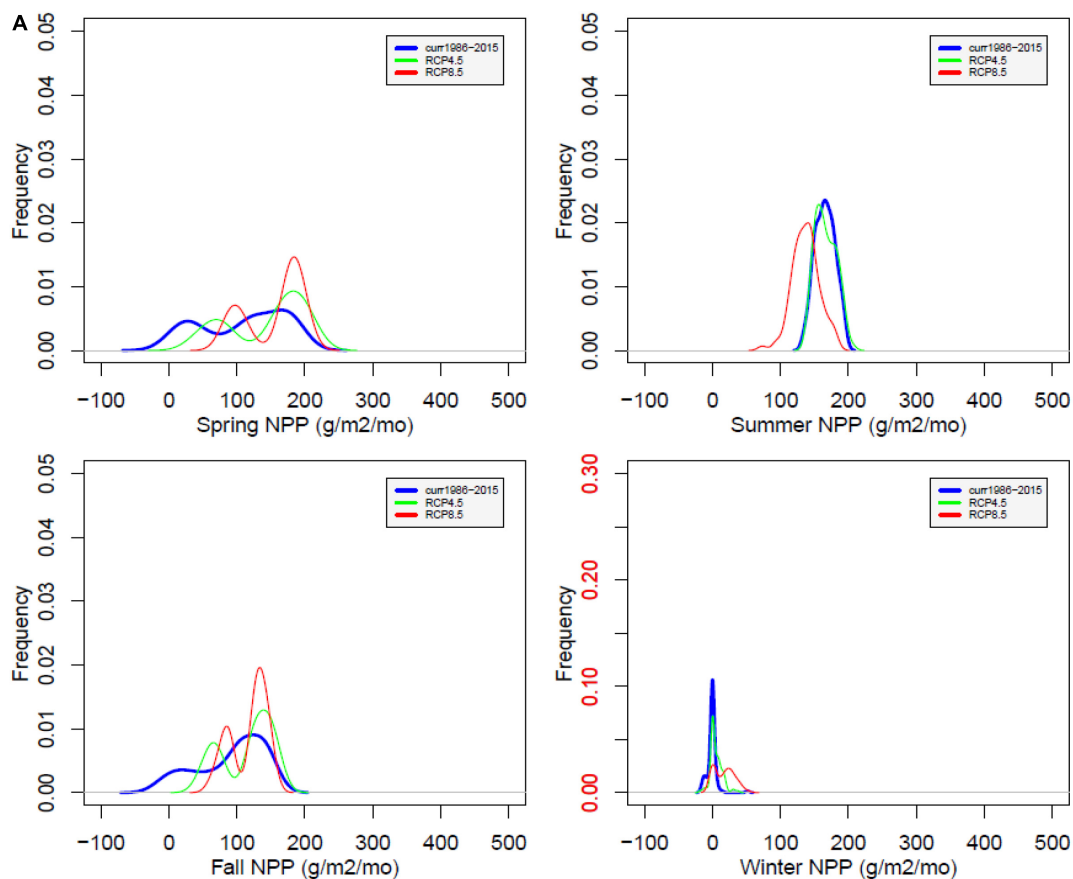


FIGURE 6 | (Continued)

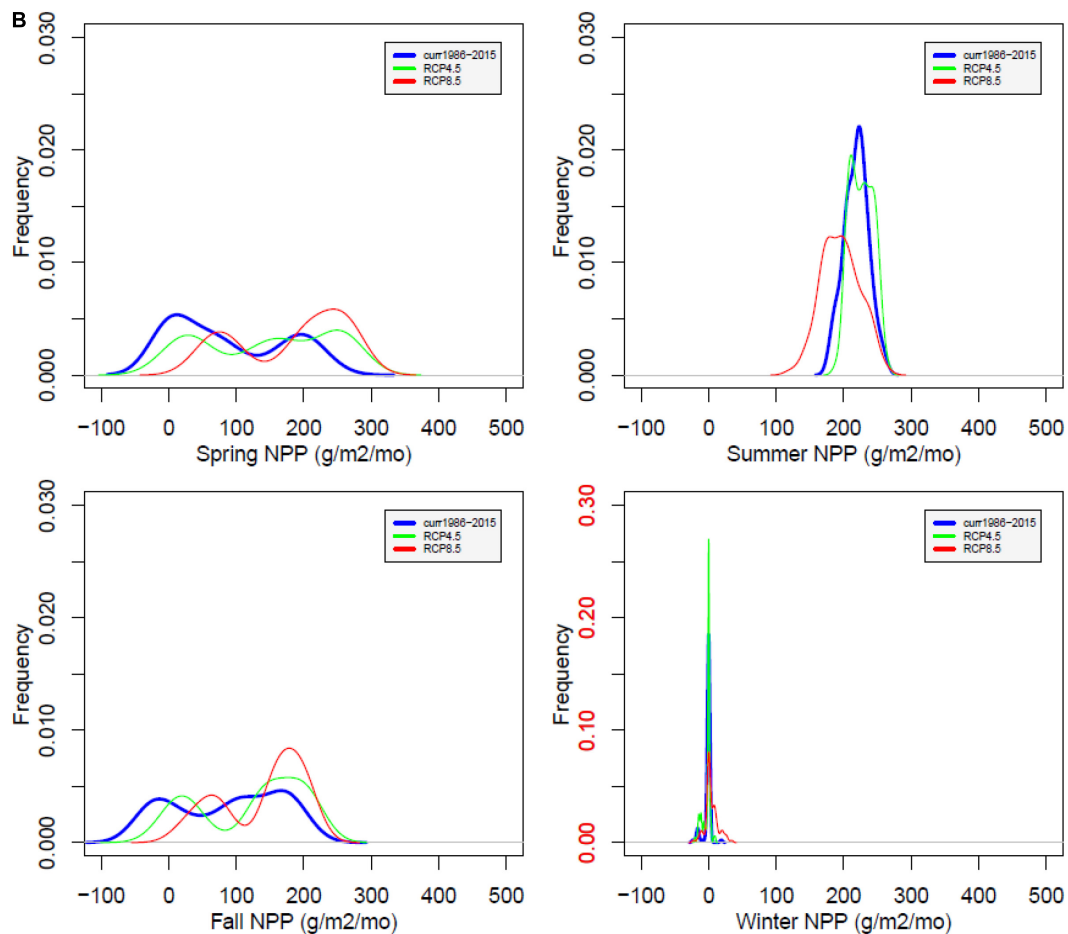


FIGURE 6 | Probability distribution of simulated net primary productivity (NPP) at WS18 (A) and WS27 (B) under future climate compared to values under current climate by season. Current (blue), RCP4.5 (green), and RCP8.5 (red). The red color of the labels in winter shows different range of y-axis compared to the other seasons.

(Supplementary Figures 1–7). As transpiration, GPP, and NPP are negligible in winter, the probability density distributions are very concentrated, unlike in other seasons.

For both watersheds, transpiration (Figure 5) is projected to increase under changing climate. However, the increase in transpiration at WS18 is smaller than at WS27, probably due to higher water stress. Even though transpiration increases, annual streamflow (Supplementary Figure 1) under future climate is projected to increase with increasing precipitation (26.8 and 32.7% at WS18 under RCP4.5 and 8.5; 11.9 and 14.7% for WS27 under RCP4.5 and 8.5, respectively). Seasonally, winter is the season that shows pronounced increase at both watersheds.

The simulated impact of climate change on GPP is not as dramatic as for the water cycle (Supplementary Figure 2). However, it shows increases in spring and fall under both climate scenarios, increases in summer under RCP 4.5, but slight decrease under RCP 8.5. The increase of GPP at WS27 is more dramatic, especially in spring, summer and fall. Overall, annual GPP is projected to increase due to the extended growing season (2071–2100 compared to 1986–2015: increase by 13.8

and 17.9% for WS18, and 26.9 and 43.0% for WS27 under RCP4.5 and 8.5 scenarios, respectively). The simulated impact of climate change on NPP is moderate (Figure 6). On an annual basis, NPP is projected to increase although decreases are evident in the summer due to increases in respiration. Under RCP8.5, NPP shows decreases in summer and a larger variance in winter, with some high values. During summer, NPP is lowered in WS18 as much as 43.2 g biomass/m²-mo or 23.1% under RCP8.5. For WS27, simulated NPP increases during all seasons under RCP4.5. With RCP8.5, NPP shows decreases during summer. NPP is projected to increase to a greater degree in WS27 than WS18 due to less water stress associated with temperature increase.

At WS18, stream ANC is projected to decrease during all seasons under future climate with the larger decrease under RCP 8.5 at WS18 (Figure 7). The impact of climate change on stream ANC is minimal at WS27, with small decreases under RCP8.5.

Under climate change, the most conservative element, Cl⁻, as well as basic cations (Na⁺, K⁺, Ca²⁺, and Mg²⁺) and SO₄²⁻ exhibited decreases in concentration but increases in

total flux associated with streamflow increases. The flux of the base cations in soil is mainly associated with release from wood litter. Calcium concentrations (**Supplementary Figure 3**) in streamwater decrease dramatically in all seasons at WS18. A similar decrease is also evident in WS27, but not as marked. Potassium concentrations decrease (**Supplementary Figure 4**) in stream water during all seasons at WS18 with the most distinct change occurring in winter and spring. Decreases in potassium are also evident at WS27 in all seasons with the most distinct change occurring in winter. Sulfate concentration (**Supplementary Figure 5**) in stream water decreases in both watersheds, especially in winter under RCP8.5 and with lower values than projected under RCP4.5 at WS18. Compared to the conditions under current climate, future sulfate concentration will be less varied at WS27 but showing more distinct change than at WS18. WS18 has the highest median sulfate concentration in summer while at WS27, the highest concentrations occur in fall and winter.

Soil processes and properties are predicted to change as well, including soil base saturation, Al:Ca concentration in soil, and nitrogen mineralization rate. Soil base saturation is indicative of the soil acid-base status with lower values symptomatic of

more acidic soil. High concentrations of Al:Ca in soil can be toxic to biota.

Soil base saturation (**Supplementary Figure 6**) shows decreases during all the seasons at both watersheds under RCP4.5 and 8.5. Decreases are pronounced under both climate scenarios at WS18, but only marginally under RCP8.5 at WS27. Al:Ca in soil shows a dramatic increase in all the seasons, but increases are most pronounced during summer at WS18 under RCP4.5 and 8.5. Al:Ca in soil increases particularly during summer at WS27. The simulated impact of climate change on N mineralization is small (**Supplementary Figure 7**). In WS18 under RCP8.5, the N mineralization rate shows some decrease. At WS27, the mineralization shows some decrease in winter but increase in summer.

Overall, the response of individual forest processes to climate change varies by season. Furthermore, vegetation seems to be more sensitive to climate change at WS27 than WS18, while soil processes and stream chemistry seem to be more impacted by climate change at WS18 than at WS27.

To more comprehensively present the impact of climate change, we summarized the changes of the 17 model-derived

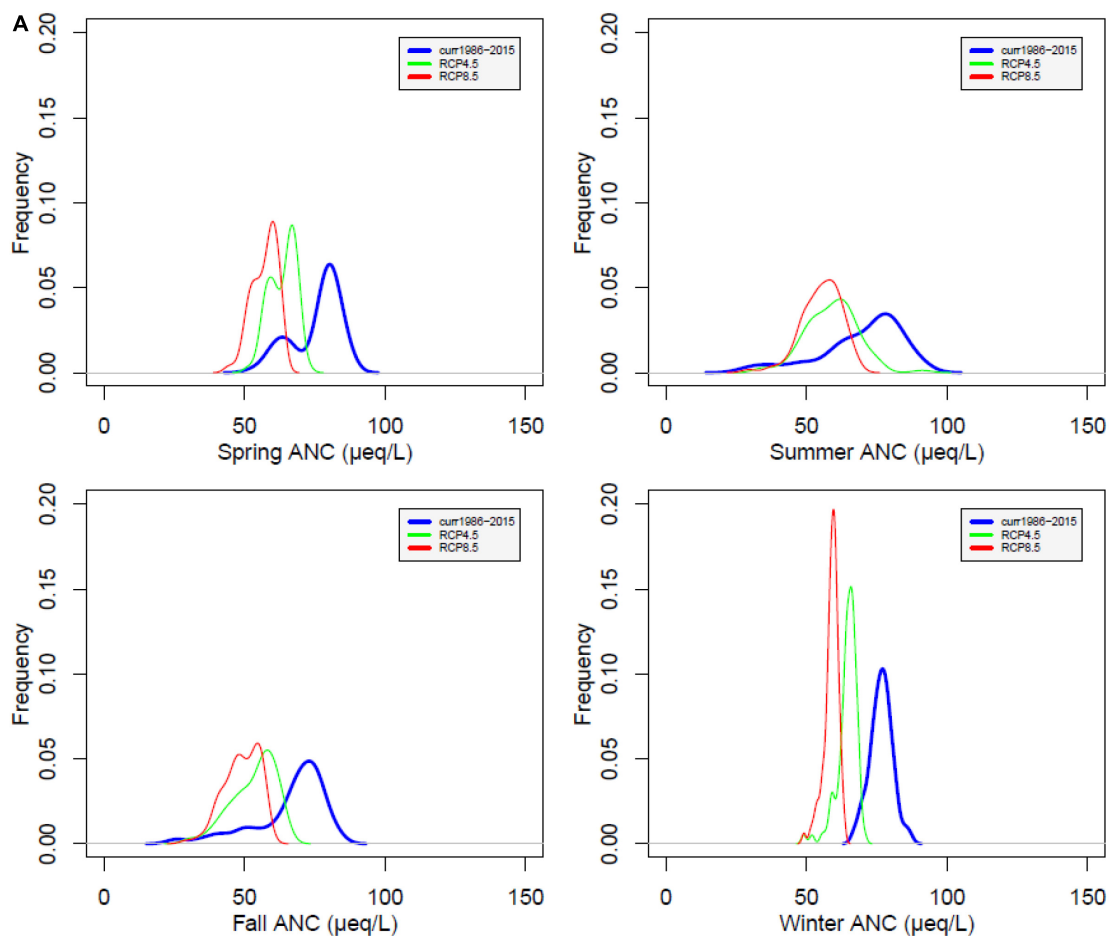


FIGURE 7 | (Continued)

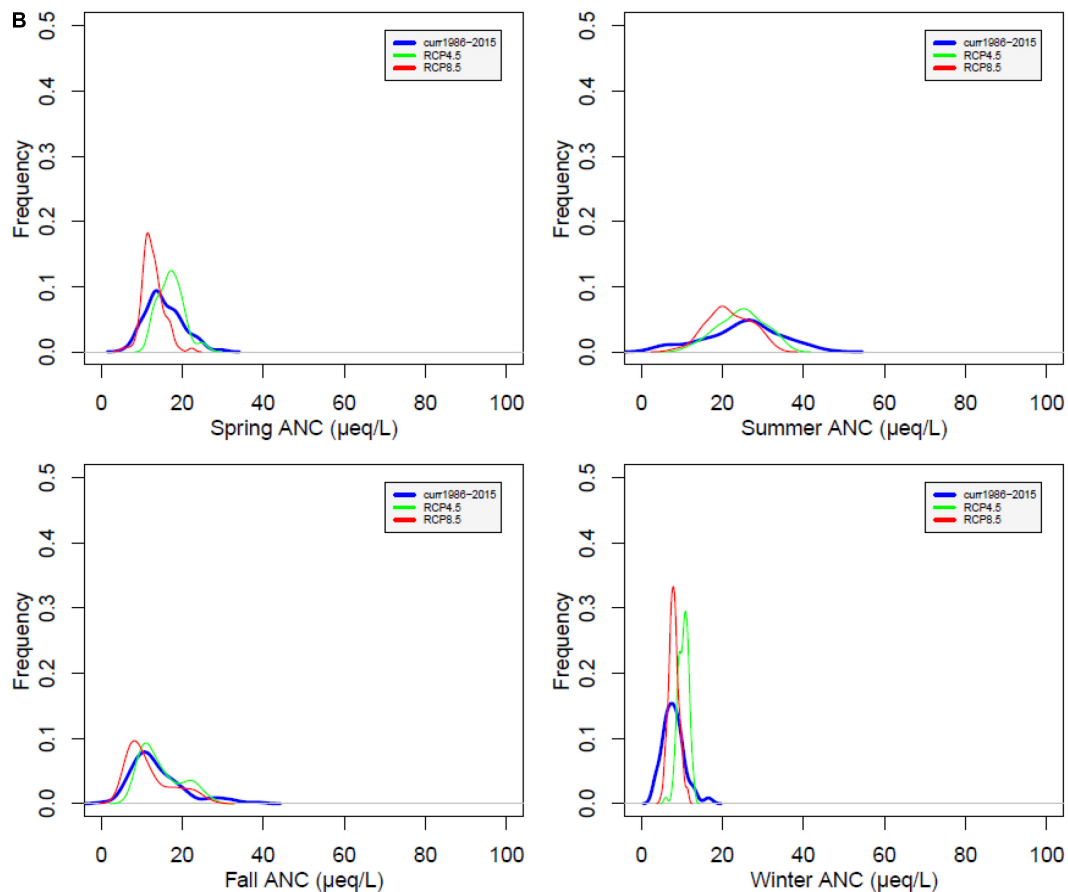


FIGURE 7 | Probability distribution of simulated acid neutralizing capacity (ANC) at WS18 (A) and WS27 (B) under future climate compared to values under current climate by season. Current (blue), RCP4.5 (green), and RCP8.5 (red).

key watershed process variables in each season between current (1986–2015) and future (2071–2100) for both RCP4.5 and 8.5 scenarios, particularly, the directions of the changes and the significance levels of the statistical tests (Table 3).

In the vegetation processes, GPP and NPP are predicted to increase significantly except summer under RCP8.5. This can be explained by higher water stress during this season.

In the hydrology process, transpiration will significantly increase during almost all seasons as both precipitation and temperature are projected to increase from present day to the end of this century. Streamflow at the Coweeta basin is trending to increase in winter and spring seasons and decrease in summer and fall seasons, but the predictions in the future are not significantly different from current conditions except winter under RCP8.5.

The soil processes captured in base saturation across both watersheds and seasons are predicted to significantly decrease, which indicates a greater loss of base cations from the soil under changing climate. Along with the reduction of base saturation, the Al:Ca ratio is predicted to increase at all seasons under both climate scenarios, and the increase is statistically significant in all seasons. Additionally, some nitrogen processes

are negatively impacted, such as gross nitrogen mineralization and immobilization. For the net nitrogen mineralization, results are complicated due to counteracting processes being involved at the same time.

In streamwater chemistry, the concentrations of major cations and sulfate are predicted to significantly decrease, even though soil losses of cations to nearby water flows (streams) are predicted to increase, probably due to the higher streamflow. ANC shows significant decrease in four seasons at WS 18 under both climate change scenarios and spring, summer, and fall at WS 27 under RCP8.5. The change of nitrate concentration mostly shows decrease.

We also found broad-scale climatic variability like El Niño–Southern Oscillation (ENSO) affected seasonal biogeochemical processes. Oceanic Niño Index (ONI that indicates ENSO) had a negative relation with the integrated biogeochemical processes (PC1) in the winter and early spring (January to April) at WS18 and WS27 (Supplementary Figure 8). Strong El Niño years generally had lower PC1 (Supplementary Figures 8, 9), indicating lower productivity and higher ANC in soil and streams, probably due to lower temperature in winter–early spring.

TABLE 3 | Significance test of 17 watershed state variables between current (1986–2015) and future (2071–2100) for the two studied watersheds (GPP, Gross Primary Productivity; NPP, Net Primary Productivity; Totlitter, Total litter produced; ANC, Acid Neutralizing Capacity; Trans, Transpiration; BS, Base Saturation; GrossNMin, Gross Nitrogen mineralization rate; GrossNImm, Gross Nitrogen Immobilization rate; NetNMin, Net Nitrogen mineralization rate; Nupdate, Nitrogen uptake rate).

	WS18								WS27							
	RCP4.5				RCP8.5				RCP4.5				RCP8.5			
	Spr	Sum	Fall	Win	Spr	Sum	Fall	Win	Spr	Sum	Fall	Win	Spr	Sum	Fall	Win
GPP	***	***	***	***	***	—	***	***	***	***	***	**	***	***	***	***
NPP	***	—	***	***	***	***	***	***	***	***	***	—	***	***	***	***
Totlitter	***	***	***	—	***	***	***	***	***	***	***	***	***	***	***	***
Mg ²⁺	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
K ⁺	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
Ca ²⁺	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
NO ₃ [−]	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
SO ₄ ^{2−}	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
ANC	***	***	***	***	***	***	***	***	**	—	—	***	***	*	*	—
Streamflow	—	—	—	—	—	—	—	*	—	—	—	—	—	—	—	**
Trans	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
Al/Ca	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
BS	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***	***
GrossNMin	—	—	—	**	***	***	***	***	—	—	—	**	—	**	—	**
GrossNImm	***	***	***	***	***	***	***	***	*	—	—	*	*	—	**	***
NetNMin	***	***	—	—	***	***	*	—	—	—	—	***	—	***	*	—
Nuptake	***	***	*	**	***	***	**	***	*	—	—	—	***	***	***	***

For significance level: *** for $p < 0.05$ and > 0.01 ; ** for $p < 0.01$ and > 0.001 ; **** for $p < 0.001$; and '—' for $p > 0.05$. Green color indicates an increase and red color indicates a decrease from current to the future climate scenarios.

Temporal Trend of Biogeochemical Processes and Change Points

From the time series of the scores of the PC1s that explained more than 50% of the total variances, we derived five similar decadal change points for WS 18 and 27 (Table 4 and Figure 8). The change points occurred earlier at WS18 than at WS27 during 1980s and 2010s but are projected to occur at similar time points or even earlier at WS 27 relative to WS18 under future changing climate. This shift may imply that future climate might accelerate changes of biogeochemical processes at the higher-elevation WS27 site. The last change points are predicted to occur 20 years earlier under RCP8.5 (2059 at WS18 and WS27) compared to those under RCP4.5 (2079 at WS18 and 2074 at WS27) at the lower- and higher-elevation forests, respectively. The groups based on the time of change points of the PC1s confirmed their validity due to minimum overlaps between these different groups (Figures 3, 4).

When we compared the change points of temperature and precipitation to those of PC1, we found the first few change points of PC1 matched those of temperature at WS18, implying temperature change is likely an important driver of watershed response (Table 4 and Supplementary Figure 10). However, precipitation change appears to be the main driver for the change points of PC1 in 2079 at WS18. At WS27, precipitation is likely the main driver for the change points in 2010s and the 2070s, while temperature is the main driver for the change points in 1989, 2030s and around 2060. The climatic driver is not clear for the change point in 1969. In general, the last change points of

PC1 under RCP4.5 (2070s) at both watersheds are likely driven by precipitation, while they seem to be driven by temperature under RCP8.5 (2050s–2060s).

DISCUSSION

Spatial and Seasonal Variability

In this analysis, we found vegetation is affected by climate change more at the higher elevation than at lower elevation, consistent with Lamprecht et al. (2018). The impacts of climate change on the hydrochemical processes have shown large spatial variability. Evapotranspiration and streamflow are predicted to increase under both moderate (RCP4.5) and high-end (RCP8.5) climate change scenarios at both WS18 and WS27. The predicted increase of evapotranspiration with climate change is consistent with patterns found in seven forest watersheds across four northeastern states in the United States (Pourmokhtarian et al., 2017). Annual streamflow, however, is highly variable among these watersheds, with some watersheds predicted to increase while others predicted to decrease, depending on whether the increase of evapotranspiration is offset by the increase of precipitation (Campbell et al., 2011). The hydrological predictions are different at the Andrews Experimental Forest in the northwestern U.S., where it is predicted that there will be a 20–71% decrease in annual transpiration under future climate change corresponding to 49–86% decreases in foliar biomass due to drought stress (Dong et al., 2019). The decrease in

TABLE 4 | Change points of the scores of the first principal components (PC1s) and climate (temperature and precipitation).

		CP1	CP2	CP3	CP4	CP5
WS18	PC1-RCP4.5	—	1981	2010	—	2039
	PC1-RCP8.5	—	1985	2010	2023	2039
	Temperature-RCP4.5	—	1984	—	2023	—
	Temperature-RCP8.5	—	—	2010	—	2038
	Precipitation-RCP4.5	—	—	—	—	2076
	Precipitation-RCP8.5	—	—	—	2043	—
WS27	PC1-RCP4.5	1969	1989	2015	—	2038
	PC1-RCP8.5	1969	1989	2017	—	2059
	Temperature-RCP4.5	—	1989	—	2038	—
	Temperature-RCP8.5	—	1997	—	2038	2067
	Precipitation-RCP4.5	—	1998	2012	—	—
	Precipitation-RCP8.5	—	1998	2012	—	—

The results are average of four climate models for two future climate scenarios.

evapotranspiration could contribute to higher streamflow at this forest. In the western U.S. where future climate is projected to become warmer and dryer, gross and net nitrogen mineralization are predicted to increase, and NPP is also predicted to increase, due to feedbacks between warmer temperatures and enhanced nitrogen mineralization in forests. In the Coweeta Basin, Knoepp and Swank (1998, 2002) found that temperature and temperature-moisture interactions significantly affected net soil nitrogen mineralization based on a series of long-term studied plots at different elevations. When temperature was warmer and soil moisture content was higher, nitrogen mineralization was reported to be greater, consistent with our findings (Table 3). From the relatively low elevation oak-pine dominated watershed to the higher elevation watershed dominated by northern hardwoods, net nitrogen mineralization rate showed a strong spatial pattern with the highest rate observed at the highest elevation site, even though other factors (e.g., C:N ratio in soil) besides temperature and soil moisture influenced soil nitrogen processes both in the laboratory and *in situ*.

Climate change is projected to have variable effects on ecosystem responses across the seasons. Even though annual streamflow is projected to increase, these increases are less pronounced in the growing season. Spring may be impacted substantially because of loss of snowpack (Reinmann et al., 2018) and early leaf-out timing that can extend the growing season and exacerbate drought stress. A hotter summer together with a warmer spring, coupled with temporally unevenly distributed precipitation, could increase drought stress and risks of wildfire (Table 3). The drought stress could extend to other seasons. The fall season presents the lowest percent increase of precipitation among all the seasons. With higher evapotranspiration driven by higher temperature, streamflow is projected to decrease thereby causing more droughts. During winter, relative temperature (three times greater at WS27 in winter under RCP8.5) increases are particularly pronounced, which decreases precipitation inputs as snow and the magnitude and duration of snow coverage (Table 1A, Campbell et al., 2011; Pourmokhtarian et al., 2017).

Effects of climate change are difficult to evaluate during the “shoulder seasons” due to the large year to year variability. As a result, impacts during these periods are generally ignored, however, more and more research shows transitional seasons are affected by climate change to a greater degree than previously thought (Xie et al., 2015; Goss et al., 2020). Traditional definition of seasons and transition between seasons are predicted to become less clear as we may experience shorter spring and fall, while winter and summer will become longer, and most of the year will fall into more extreme hot and cold temperatures (Thomson, 2009). The impact of climate change on biogeochemical process during the shoulder transitions needs to be studied further.

The Impact of CO₂ on Ecosystems

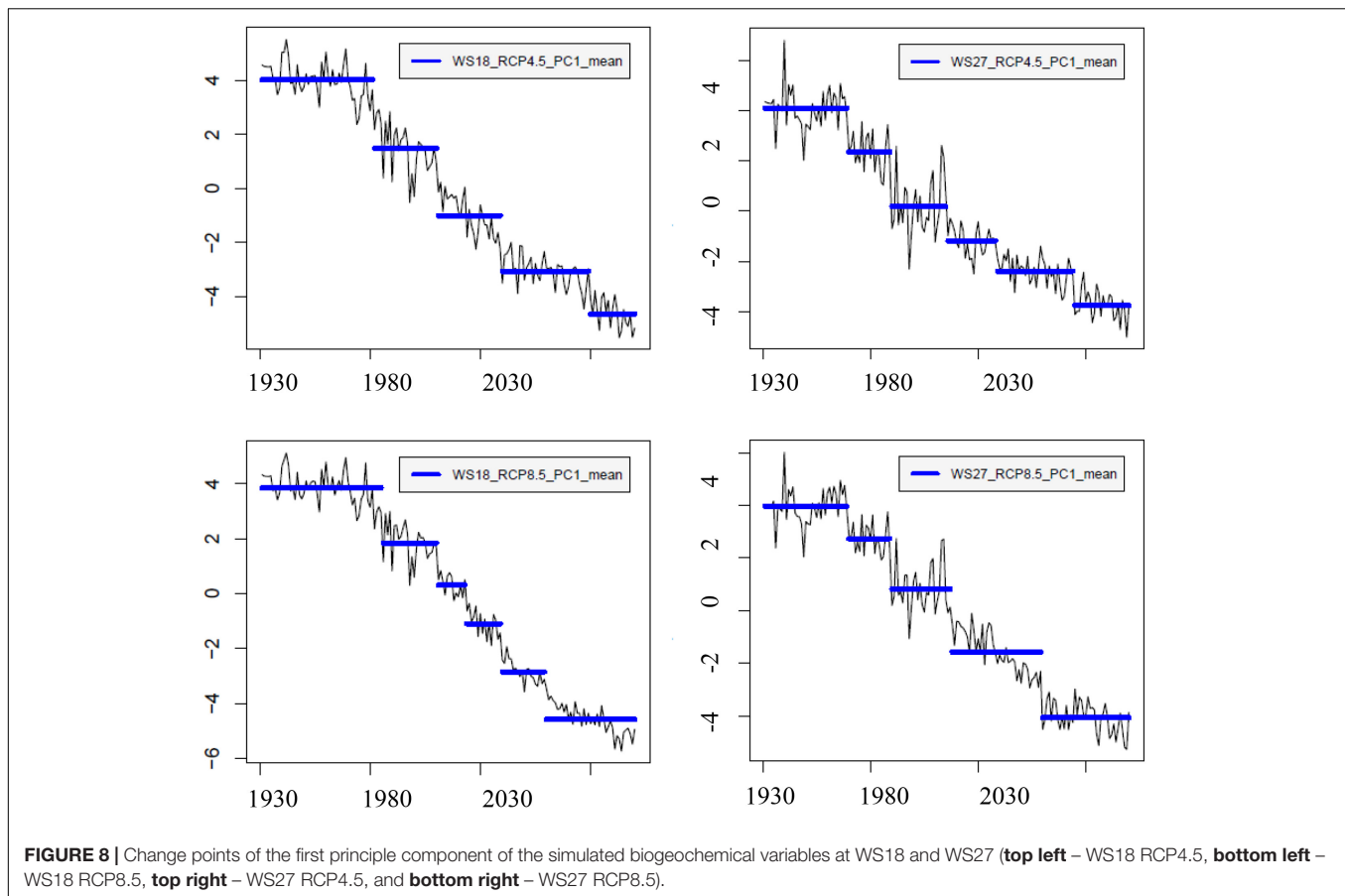
Elevated CO₂ associated with climate change tends to increase plant WUE, i.e., increases primary productivity and reduces evapotranspiration (Reyer et al., 2015), which benefits forests by increasing their drought tolerance although there are counteracting effects to consider. Elevated CO₂ could also increase plant leaf area, therefore, making trees less drought tolerant (Ghannoum and Way, 2011). Reduced evapotranspiration could also lead to increases in leaf temperature, thereby increasing temperature stress. Increased productivity under elevated CO₂ may increase litter (Hyvonen et al., 2007), leading to increased litter accumulation (if decomposition cannot catch up) and higher vulnerability to fire. Many studies on the response of forest habitats to increased CO₂ are made at the small scale of individual trees or small plots. Scaling up the effect to the ecosystem scales involves considerable uncertainty. Furthermore, the fertilization effect of CO₂ is commonly found only in young trees (Korner et al., 2007). In our study, we assume elevated CO₂ increases primary productivity at the entire watershed without considering other counteracting effects (e.g., leaf temperature increase or age of trees), therefore our prediction of primary productivity under climate change is likely an optimistic estimate.

Uncertainties of Model Predictions

It is important to predict climate as accurate as possible in order to make valid predictions of ecosystem function change. This has been facilitated using downscaling method in our study. However, broad-scale climate variability like ENSO may enhance changes in temperature or precipitation (El Niño and La Niña Years and intensities at⁸ and accessed on 08/01/2020) at local spatial scales, and therefore has impact on biogeochemical processes as what we found out in this study. Generally, during El Niño years, the southeastern U.S. experiences cooler temperatures and wetter conditions in winter-early spring⁹ accessed on 08/01/2020. Together with the North Atlantic Oscillation (NAO), it was found that the southern Appalachians have higher snowfall and cooler temperature in winter during El Niño and negative NAO phases (Eck et al., 2019). In contrast, La Niña and positive NAO years result in warmer and drier winter

⁸<https://ggweather.com/enso/oni.htm>

⁹<https://www.weather.gov/tae/enso>



weather. As the prediction of El Niño's intensity using current climate models involves large uncertainty (Wang et al., 2019), predictions of temperature, precipitation and biogeochemical processes are particularly challenging under climate change. In order to reduce uncertainties of predicted biogeochemical processes, broad-scale climate variability needs to be accounted for in the climate change models.

PnET-BGC is a watershed-scale model that is operated on a monthly time step, so it is not well suited to address finer spatial and temporal resolutions (Merganicova et al., 2019). This model treats the vegetation in the watershed as a big leaf, therefore, it does not consider the difference among tree types and fine structures (Elliott et al., 2015), adaption (Jandi et al., 2019), or explicitly accounts for shift of vegetation types. Previous studies show that the critical transitions of vegetation type in mountain forests (Reich et al., 1995; Albrich et al., 2020) may change the fundamental relationships between foliar N concentration and maximum photosynthesis rate, which is one of the two fundamental relations that PnET-BGC is built on (Gbondo-Tugbawa et al., 2001). Furthermore, the function of understory shrub vegetation in cycling nutrients, carbon, and water (Rothstein, 2000) is another component not considered in this model. Different stages of vegetation not only have different capabilities of nutrient uptake, photosynthesis, cycling of carbon and water, but larger and older trees may be also more susceptible to stress such as drought and storms (Clinton

et al., 1993, 2003; Clinton and Baker, 2000), which is not depicted in the model.

This PnET-BGC model requires a number of empirical data to calibrate, some of which are not available for the study watersheds (such as Al and DOC concentrations), making calibration for these parameters less robust. In addition, the model under-predicts nitrate concentrations in streamwater, suggesting the need to critically review this submodule for applications at Coweeta watersheds. Other simulations using PnET-BGC have been challenged to accurately simulate nitrogen dynamics (e.g., Shao et al., 2020). This model is designed to predict long-term hydrologic and biogeochemical function at the watershed scale (Aber and Federer, 1992; Gbondo-Tugbawa et al., 2001), combined with the lack of feedback between vegetation and climate, it is not an ideal tool to derive regime shifts. Other components lacking in the PnET-BGC further contribute to uncertainties in the simulations. These include the lack of multiple soil depths (two-layer model available but harder to calibrate so we used one-layer model), and lack of consideration of phosphorus dynamics.

Despite these aspects that can be improved, PnET-BGC provides a useful tool to evaluate the impact of climate change on forested ecosystems over long time scales (multiple decades to centuries). When depicting complex ecosystems, as many other ecological models have done, it is necessary to make

assumptions and acknowledge inherent uncertainties (Aber and Federer, 1992). In order to capture the uncertainty, we have applied two climate change scenarios. The consistent response of biogeochemical cycles provides higher confidence in our predictions on the climate change impact.

Implications for Forest Management

This research will provide useful information for resource managers in anticipation of the potential risks forests may face under climate change. Forest managers can consider possible measures such as tree planting and forest restoration, wildfire control, establishment of mixed stands, planting better adapted species or varieties to mitigate adverse impacts in the future. For monitoring, the forest managers may consider watershed specific and seasonal specific evaluations instead of focusing on growing seasons or one process in soil or vegetation only.

We are facing an uncertain future based on (1) the greater uncertainty in prediction for precipitation than for temperature (Intergovernmental Panel on Climate Change [IPCC], 2014), and (2) precipitation will be the main driver for future change points under RCP4.5. Considering the change that may occur in high-elevation forests directly influences ecosystems downslope and downstream and the highly uncertain future projections, measures should be taken to protect, conserve and restore these high-elevation forests. The improved understanding of forests' non-linear responses to a changing environment gained from this research can potentially guide decisions in forest management even with large uncertainties, for example, detection of early warning signs of potential regime shifts and selection of species planted in restoration.

CONCLUSION

We predict both positive and negative impacts of climate change on hydrochemical processes, for example, we predict increases in acidification of soil and streamwater despite an increase in primary productivity under climate change. The predicted hydrochemical changes show strong seasonal variability depending on biogeochemical processes. We also find climate change will likely have a greater impact on soil processes at the lower elevation watershed and on vegetation processes at the higher elevation watershed. The change-point analysis suggests biogeochemical cycles in vegetation, soil, and streams may be accelerating at higher-elevation. These findings that include integrated hydrochemical processes in temporal and spatial (lower-elevation vs. higher-elevation) contexts provide multi-dimensional metrics that can be used in part or as a whole by southern forest managers to evaluate the benefits

and disadvantages in anticipation of future climatic change to improve the resilience of forest ecosystems and downstream surface water in the future. Additionally, the integrated modeling framework can be applied to specific management practice scenarios of interest to yield more detailed quantitative predictions for in-depth analysis of these management plans.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/**Supplementary Material**, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

KE and CM provided measured data at the Coweeta Basin. HH performed the PnET-BGC model runs with the guidance of WW and CD. HH and WW interpreted the model results and wrote the first draft of the manuscript. All authors commented on previous versions, and read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/ffgc.2022.853729/full#supplementary-material>

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